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(71) Applicant (for all designated States except US): THE WELLESLEY HOSPITAL FOUNDATION [CA/CA]; 160 Wellesley Street East, Toronto, Ontario M4Y 1J3 (CA).

(72) Inventors; and

- (75) Inventors/Applicants (for US only): JULIUS, Michael, H. [CA/CA]; 38 Earl Street, Townhouse #9, Toronto, Ontario M4Y 1M3 (CA). FILIPP, Dominik [SK/CA]; 704 15 Dundonald Street, Toronto, Ontario M5Y 1K4 (CA). ALIZADEH-KHIAVI, Kamel [IR/CA]; 618 1201 Richmond Street, London, Ontario N6A 3L6 (CA).
- (74) Agent: HUNT, John, C.; Blake, Cassels & Graydon, Box 25, Commerce Court West, Toronto, Ontario M5L 1A9 (CA).

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#### (57) Abstract

A novel protein purified from bovine colostral whey and isolated nucleotide sequences encoding the protein is identified. The isolated bovine protein is termed Bovine Lactation Associated Immunotropic (Bo-LAIT) protein. The human homologue of Bo-LAIT protein, Hu-LAIT protein, is also described. A method of activating B cells, and particularly of activating B cells in a mammal, such as a human, in need of such activation by administering LAIT protein is described. LAIT protein can be incorporated into infant formula. LAIT protein can be administered to an infant, as by feeding to the infant such formula. LAIT protein can be incorporated as part of a vaccination. LAIT protein can be administered to a patient having a T cell immune deficiency, for example, a particular T cell dysfunction in which gp39 (CD40L) is under expressed on or totally absent from the cell surface of patient T cells. Preparation of medicaments including LAIT protein for activating B cells in a mammal in need of such activation is described. Natural or recombinant LAIT protein can be used.

BOVINE LACTATION ASSOCIATED IMMUNOTROPIC PROTEIN (CD14), ENCODING GENE AND APPLICATION IN B CELL ACTIVATION

#### FIELD OF INVENTION

This invention, in the fields of immunology, biochemistry and cell and molecular biology, relates to proteins or proteins that are co- and/or post-translationally modified, termed LAIT proteins, that activate B cells. This invention is also directed to the use of such protein in pharmaceutical preparations, and pharmaceutical compositions comprising LAIT protein or functional derivatives thereof. This invention is also directed to nucleic acid molecules encoding the bovine LAIT protein or functional derivatives thereof and methods for the purification of native and recombinant forms of said proteins that activate B cells.

### **BACKGROUND OF THE INVENTION**

Bone marrow-derived "B" lymphocytes, commonly called B cells, are a type of white blood cell present in the lymph, the blood, and in secondary lymphoid organs of the immune system. B cells are the precursors of antibody secreting cells, plasma cells, and as such are central to the induction of humoral immune responses.

The induction of most humoral immune responses in the adult involves a number of cellular interactions among thymus-derived T lymphocytes, commonly called T cells, antigen presenting cells (APC), and B cells [J. Exp. Med 147:1159, 1978; PNAS 77:1612,1982; PNAS 79:1989, 1982; Immunol. Rev. 95:914, 1987].

As currently understood, T cell-dependent B cell activation involves activation of T cells upon their recognition of antigen, as presented by APC in conjunction with proteins encoded within the major histocompatibility complex (MHC), which are expressed on the cell surface of the APC. This antigen specific and MHC restricted T cell-APC interaction results in reciprocal activation of the two cell types, and the alteration of T cell physiology such that "helper function" becomes manifest.

Helper T cells can activate antigen specific B cells. Antigen specificity of the T cell-B cell interaction is maintained as a consequence of the ultimate capacity of the B cell to function as an APC. Thus, while resting, quiescent B cells are not efficient APC (PNAS 79: 1989, 1982), they specifically interact with antigen through membrane associated immunoglobulin, the specificity of which reflects that of the immunoglobulin their daughter cells will secrete (J. Exp. Med. 140:904, 1974).

Immunoglobulin mediated internalization of antigen by the specific B cell, which may involve presentation by yet another sort of APC, the follicular dendritic cell, results in the initiation of antigen processing by the B cell, the up-regulation of MHC Class II and B7 expression, and the presentation of antigen derived peptides in the context of MHC (J. Exp.

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Med. 178: 2055, 1993). The B cell activated by this route is a target for the activated helper T cell.

T cell helper function includes signals delivered through both T cell-B cell contact, and the interaction of T cell derived soluble mediators, referred to as cytokines, with their cognate ligands expressed on the B cell plasma membrane. T cell-B cell contact is also MHC restricted, analogous to the T cell-APC interaction (Eur. J. Immunol. 12:627, 1982; Eur. J. Immunol. 12:634, 1982). However, the specific interaction of the molecules which mediate the MHC restricted interaction between the two lymphocyte lineages, specifically, the T cell receptor for antigen (TcR), and the MHC/antigen complex expressed by the B cells, do not predicate the induction of B cell growth and differentiation (Eur. J. Immunol. 18:375, 1988).

The essential molecular interaction, reflected by the requirement for T cell-B cell contact, is mediated by CD40 expressed on the plasma membrane of the B cell, and its cognate ligand, gp39 (or CD40L), expressed on the plasma membrane of the T cell (PNAS 89:6550, 1992; Nature 357:80, 1992). Consistent with this paradigm is the observation that membrane expression of the latter increases upon T cell-APC interaction, as well as subsequent to T cell-B cell interaction (PNAS 89:6550, 1992). Further, membrane immunoglobulin mediated B cell interaction with antigen results in the increased membrane expression of CD40 (Sem. in Immunol 6:303, 1994). The interaction between CD40 and CD40L predicates the induction of B cell growth, B cell differentiation into immunoglobulin secreting cells, and immunoglobulin isotype switching (J. Exp. Med. 178:1567, 1993).

Consistent with this model is the observation that soluble CD40L, or monoclonal antibody (mAb) specific for CD40 can induce B cell growth and differentiation to immunoglobulin secretion (Sem. in Immunol. 6:267, 1994; PNAS 83:4494, 1986; J. Immunol. 140:1425, 1988; ).

In addition to the obligate requirement for T cell-B cell contact, a number of T cell derived cytokines, IL-2, IL-4 and IL-5 are central to B cell growth and differentiation. B cell susceptibility to these cytokines is for the most part limited by prior contact with a T cell. Thus, subsequent to T cell contact, the B cells increase expression of cytokine specific membrane receptors (PNAS 80:6628, 1983; J. Immunol. 145:2025, 1990; J. Immunol. 146:1118, 1991). IL-2 and IL-5 have been demonstrated to support the growth of activated B cells (PNAS 77:1612, 1980; Immunol. Rev. 52;115, 1980). Further, IL-4 and anti-immunoglobulin have been shown to synergize in supporting B cell growth (J. Exp. Med. 155:914, 1982).

Notable exceptions in this context are the quiescent B cell responses to IL-4 and IL-5. IL-4 induces the de novo transcription and translation of MHC Class II proteins (J.

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Exp. Med. 155:914, 1982; PNAS 81:6149, 1984; J. Exp. Med. 160:679, 1984), and IL-5 is able to support the differentiation of quiescent 8 cells into high rate immunoglobulin secreting cells in the absence of cell growth (Eur. J. Immunol. 22:2323, 1992).

In any event, signals derived from molecular interactions amongst membrane molecules on T cells and B cells, and from those of T cell derived cytokines interacting with their cognate receptors on B cells are parts of a complex signaling system. Each signal drives the B cell to another stage of activation, rendering it susceptible to subsequent progression signals. These signals complement one another, rather than having the capacity, individually, to drive the complete process of B cell growth and differentiation (Immunol. Rev. 95:177, 1987).

In 1988, a unique activity in ovine colostrum was discovered (J. Immunol. 140:1366, 1988). Proline Rich Protein (PRP) had been partially purified using classical techniques of protein purification. This material was shown to support the induction of quiescent B cells into the cell cycle, and to support their differentiation into high rate immunoglobulin secreting cells. This was apparently the first report of a protein of mammalian origin that mediates these functions.

A monoclonal antibody specific for ovine PRP was subsequently prepared. When PRP preparations were passed over an affinity column prepared using the antibody, all of the PRP was retained by the column, as assessed by Western blotting analysis of eluate and effluent. However, all of the B cell stimulatory activity was found in the effluent. Thus, the published characterization of the B cell tropic bioactivity present in ovine colostrum was not attributable to PRP (unpublished information).

### SUMMARY OF THE INVENTION

This invention features a novel bovine protein and isolated nucleotide sequences encoding the protein, the said protein being capable of activating B-cells of mammalian origin. A substantially pure LAIT protein or co- and/or post-translationally modified form of the protein may be produced by biochemical purification, or by recombinant means in a prokaryotic or eukaryotic host substantially free of other proteins with which it is natively associated. Also included in this invention is a process for purifying LAIT protein or a co- and/or post-translationally modified form of LAIT protein of this invention from bovine colostral whey comprising:

- (i) salting out of proteins contained within said samples
- (ii) enrichment and ultimate purification of LAIT protein from proteins salted out in step (i) utilizing classical protein fractionation techniques.

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In all cases the said protein possesses the desired biological activity.

The invention is also directed to a nucleic acid molecule comprising a nucleotide sequence encoding a LAIT protein. The nucleic acid molecule may be cDNA or genomic DNA.

The isolated bovine protein has homology with human CD14 and murine CD14 and so is also referred to as bovine CD14. The invention includes a method of activating B cells, and particularly of activating B cells in a mammal in need of such activation by administering CD14, a recombinant form of the protein thereof, or a functional derivative thereof.

In a preferred embodiment, the mammal is a human patient.

According to one aspect, the invention includes incorporating CD14 into infant formula. The invention includes administering CD14 to an infant, a preferred mode of administration being feeding to the infant such formula.

In another aspect, the invention includes incorporating CD14 as part of a vaccination. The invention includes administering CD14 and antigen to a patient in need of immunization, a preferred mode of administration including administering a single preparation containing both CD14 and the antigen.

In another aspect, the invention includes administering CD14 to a patient having a T cell immune deficiency. In a preferred aspect, the invention includes administering CD14 to a patient suffering from a particular T cell dysfunction in which gp39 (CD40L) is under expressed on or totally absent from the cell surface of patient T cells.

In another aspect, the invention includes administering antibodies raised against CD14 to a patient suffering from a dysfunction wherein the patient's B cells are hyperactivated as a result of higher than normal levels of serum CD14. In a preferred aspect, the invention includes administering antibodies against CD14 to a patient suffering from rheumatoid arthritis wherein the B cells are secreting rheumatoid factor as a result of being activated by serum CD14.

This invention includes a novel method of the production of hybridomas secreting mAb of desired specificity by cuturing B cells with sub-optimal mitogenic concentrations of CD14 in concert with the antigen to which antibodies wished to be raised against. Populations of B cells activated in this manner are highly enriched for activated, antigen specific B cells, which are then be used for the production of hybridomas secreting the mAb of desired specificity.

The invention includes use of CD14 in preparation of medicaments for activating B cells in a mammal in need of such activation.

Natural or recombinant CD14 can be used in the invention.

In the context of this invention, the term "CD14" includes murine, bovine or human CD14.

### **DEFINITION OF TERMS**

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A "functional derivative" retains at least a portion of the function of CD14, such as binding to a specific antibody or binding to its cognate receptor on cells that possess said receptor which permits its utility in accordance with the present invention. The term "functional derivative" as used herein includes a "fragment," or "variant" of CD14, which terms are defined below.

A "fragment" of CD14 refers to any subset of the polypeptide, that is, a shorter peptide. The term "fragment" is used to indicate a polypeptide which is derived from CD14 having a naturally occurring protein sequence comprising a deletion of one or more amino acids at one or more sites of the C-terminal, N-terminal, and within the sequence. Such fragments should retain one or more biological activities or functions which are characteristic for the intact CD14 polypeptide or co- and/or post-translationally modified forms of CD14.

A "variant" of CD14 refers to a polypeptide having a primary sequence similar to that of the native CD14 or fragment thereof such that native activity is at least partially retained. Variant peptides may be prepared by synthetic means or by mutations in the cDNA encoding said polypeptide that retains biological activity of said polypeptide including deletions, insertions or conservative amino acid substitutions within the polypeptide.

The term "antibody" as used herein is an immunoglobulin protein that has the capability to bind a distinct epitope in an unconserved region of said protein thereby enabling the antibody to distinguish one protein from another. The term "epitope" is meant to refer to that portion of any molecule capable of being bound by an antibody. The term "antibody "includes polyclonal antibodies, monoclonal antibodies (mAbs) or chimeric antibodies.

Polyclonal antibodies are heterogenous populations of antibody molecules derived from the sera of animals immunized with an antigen.

Monoclonal antibodies are a homogenous population of antibodies capable of binding a distinct epitope on the antigen. MAbs may be obtained by methods known to those skilled in the art. Such antibodies may be of any immunoglobulin class including IgG, IgM, IgE, IgA, and IgD, and any subclass thereof. The term "antibody" is also meant to include both

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intact molecules as well as fragments th reof, such as, for example Fab and F(ab')<sub>2</sub>, which are capable of binding antigen. Fab and F(ab')<sub>2</sub>, lack the Fc fragment of the intact antibody, clear more rapidly from the circulation, and may have less non-specific tissue binding than an intact antibody (Wahl et al., J. Nucl. Med. 24:316-325, 1983).

Chimeric antibodies are molecules, different portions of which are derived from different animal species, such as those having variable region derived from a murine mAb and a constant region derived from a human immunoglobulin.

An "antigen" is a molecule or a portion of a molecule capable of being bound by an antibody which is additionally capable of inducing an animal to produce an antibody capable of binding to an epitope of that antigen. An antigen may have one, or more than one epitope. When an antibody is said to be "specific for" a polypeptide, fragment, or variant thereof or is said to be "capable of binding" to a polypeptide, fragment, or variant thereof it is meant that the antigen will react in a highly selective manner, with its corresponding antibody and not with the multitude of other antibodies which may be evoked by other antigens.

### 15 BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1A -1C show purification of native bovine LAIT (nBo-LAIT) protein. Figure 1A shows an elution profile from an anion exchange column [FPLC-Mono-Q, Pharmacia] onto which was loaded 50 mg of the 62% (v/v) (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> precipitate. Bound proteins were eluted with a gradient of 50-400mM NaCl in 10mM Bis-tris propane, and a simultaneous pH gradient of 7.5 to 9.5. Table 1A indicates the bioactivity of the fraction with peak activity, fraction #57. High buoyant density mouse splenic B cells were isolated as previously described (J. Immunol. 131:581, 1983), and cultured at 5 x 104 cells in 0.2 ml of serum free medium in a 96 well cluster, flat bottomed tissue culture plate (Costar). Each fraction was added to a final concentration of 10% (v/v) in the presence, or absence of 0.25  $\mu$ g/ml LPS. At 40 hours, cultures were pulsed with 1 µCi of 3H-TdR, harvested onto filter discs 6 hours later, and thymidine uptake assessed by scintillation spectroscopy. Numbers represent cpm x 10<sup>-3</sup>. Figure 1B shows a profile from a molecular sieving column [FPLC-Superdex 75, Pharmacia] onto which was loaded 20 mg of Fraction #57 (Figure 1A). The column was equilibriated in 20 mM tris-HCL pH 8.0 buffer containing 0.45M NaCl. Table 1B indicates the bioactivity of the peak fraction, fraction #38, assessed as described in connection with Figure 1A. Figure 1C shows the elution profile of a hydroxy apatite column [HPLC-hydroxy apatite, Pharmacia] onto which was loaded 1mg of fraction #38 (Figure 1B). Bound proteins were eluted with a a gradient of 1-500mM K, HPO4 buffer, pH 6.8 containing 1 mM NaCl. Table 1C indicates the bioactivity of the peak fraction, fraction #25, assessed as described in connection with Figure 1A. The inset

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in the figure shows a silver stained SDS-PAGE gel of roughly 5  $\mu$ g of protein from fraction #25. Left lane: fraction #25; right lane: MW markers, from the top: 97,66,45,31,21, and 14kD, respectively.

Figure 2 shows the known sequence of human CD14 (SEQ ID NO:5) and aligned fragments of nBo-LAIT. Bo-LAIT fragments were generated from affinity purified colostral nBo-LAIT (see Figure 3). Fragments corresponding to residues 235-264 and 344-355 of human CD14 were major and minor peptides, respectively, each approximately 18kD in size, generated by CnBr cleavage, and separated by reverse phase HPLC (C8 column, Pharmacia). The fragment corresponding to residues 53-67 of human CD14 is a partial sequence of a 24kD fragment generated by CnBr cleavage, and separated by SDS-PAGE and electroblotted onto PVDF membrane. Fragments corresponding to residues 19-36 and 151-165 of human CD14 were generated by trypsin cleavage, and separated by reverse phase HPLC (C8 column, Pharmacia). The length of the overlapping bovine sequence with the predicted sequence of human CD14 is underlined for each of the fragments. Dashes indicate the same amino acids while those differing from the human sequence are indicated.

Figure 3 shows SDS-PAGE and silver staining of affinity purified colostral LAIT-protein from bovine and human. Lane 1: MW markers, as given for Figure 1C; lane 2: 62.5% (v/v) (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> precipitate of bovine colostral whey; lane 3: pH 2.5 eluate from #842-Sepharose affinity column; lane 4: pH 2.5 eluate from CD14 specific mAb 63D3 (PNAS 77:6764, 1980)-Sepharose affinity column, loaded with material represented in lane 5; lane 5: Sephacryl S100 HR fractionated human colostral whey. Each of lanes 1-5 contain 5  $\mu$ g of protein. Table 2 shows results obtained when 5 x 10<sup>4</sup> high buoyant density mouse splenic B cells were cultured in serum free medium in the presence of the indicated stimuli for 40 hours, pulsed with 1  $\mu$ Ci of <sup>3</sup>H-TdR, harvested 6 hours later onto filter discs, and thymidine uptake assessed by liquid scintillation spectroscopy. Numbers represent cpm x 10<sup>-3</sup>. Details of the bioassay are as described for Figure 1A. Control cpm x 10<sup>-3</sup>: no stimulus, 0.3; 50  $\mu$ g/ml LPS, 75.0; 0.25  $\mu$ g/ml LPS [LPS 1], 0.8; and 1  $\mu$ g/ml mlgM specific mAb b-7-6 (Eur. J. Immunol. 14:753, 1984), 0.7.

Figures 4A and 4B show heat lability and antibody mediated inhibition of nBo-LAIT activity. Figure 4A shows thymidine uptake by 5 x 10<sup>4</sup> high buoyant density mouse splenic B cells cultured as described for Figure 1A in the presence of the indicated concentration of affinity purified nBo-LAIT that had been heat treated at 95°C for 10 minutes (•), and nBo-LAIT that had not been heat treated (O). Cultures were pulsed with <sup>3</sup>H-TdR at 40 hours, harvested 6 hours later, and thymidine uptake assessed by liquid scintillation spectroscopy. The inset depicts the responses in cultures containing the indicated

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concentrations of LPS, which had been heat treated (or not) as for nBo-LAIT. Figur 4B shows thymidine uptake in cultures that were established as described for Figure 4A in the presence of either 0.25  $\mu$ g/ml of affinity purified nBo-LAIT, or 50  $\mu$ g/ml of LPS. Each of these stimuli were cultured in the presence of the indicated concentration of either polyclonal rabbit lgG anti-Bo-LAIT, #842, or normal rabbit lgG. The percent inhibition of thymidine uptake mediated by #842 lgG for both nBo-LAIT and LPS mediated stimulation is indicated in parentheses. Levels of inhibition mediated by normal rabbit lgG ranged from 9-20%, and 12-31% for nBo-LAIT and LPS stimulation, respectively. CPM directly induced by #842 lgG in isolation ranged from 454 ±53 to 764 ±69 at 0.4 and 50  $\mu$ g/ml, respectively; and for normal rabbit lgG, from 297 ±34 to 420 ±31 at 0.4 and 50  $\mu$ g/ml, respectively. Non-stimulated controls gave rise to 195 ±29 cpm for both sets of experiments.

Figure 5A shows a restriction map of the 7.1 kb EcoRl-Xhol fragment containing bovine CD14 gene. Abbreviation for restriction sites are: X, Xhol; P, Pstl; O, Ncol; B, BamHl; N, Notl; D, BssHl; T, BstEll; M, Smal; S, Sacll; C, Hpal; R, EcoRV; A, Sphl; G, Bglll; H, Hindlll; E, EcoRl. Figure 5B is a schematic diagram of the bovine CD14 locus. The shaded area represents the coding region of the gene, the open box is an intron sequence. The dashed area in front of the ATG start codon is 5' untranslated region, and the dashed area behind the TAA stop codon is 3' untranslated region. Figure 5C is a schematic diagram showing the sequencing strategy taken. Arrows represent the direction of sequencing. The fragment number is indicated at the right (see text for detail).

Figure 6 shows a comparison of nucleic acid sequences of bovine (SEQ ID NO:1), human (SEQ ID NO:2) and mouse (SEQ ID NO:3) CD14 coding regions. The first base position corresponds to the first nucleotide of the ATG codon, the last nucleotide corresponds to the third nucleotide of the TAA stop codon. Alignment was done using DNA STAR-Megalign software, applying the Clustal method with a weighted residue table. Human cDNA sequence (accession number P08571) and mouse cDNA sequence (accession number P08571) used in this alignment were derived from the Swiss-Protein Database.

Figure 7 shows a comparison of amino acid sequences of bovine (SEQ ID NO:4), human (SEQ ID NO:5) and mouse (SEQ ID NO:6) CD14 proteins. Amino acid sequences were deduced from the corresponding cDNA sequences shown in Figure 6. DNA Star-Megalign software was used to generate this alignment using the method described by J. Hein (Methods in Enzymology 183:626, 1990) in conjunction with the PAM 250 residue weight table.

Figures 8 show primers used for amplification of Bovine CD14 cDNA coding region. Figure 8A shows forward (SEQ ID NO:7) and reverse (SEQ ID NO:8) primers used for

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the baculovirus expression system. Figure 8B shows forward (SEQ ID NO: 9) and rev rse (SEQ ID NO: 10) primers used for the mammalian expression system.

Figure 9 shows immunoblotting of native and recombinant bovine CD14. Western blot analysis was used to evaluate and compare the sizes of nBo-LAIT protein with recombinant CD14 proteins. 250 ng of CD14 proteins were electrophoresed on 12.5% SDS-polyacrylamide gel and electrophoretically transferred to PVDF membrane (Millipore) at 180 mA for 30 minutes. The membrane was blocked for 1 hour in 5% skim milk in TBST (20mM Tris.HCl, pH 7.5, 150mM NaCl, 0.025% Tween 20), followed by incubation for 1 hour with rabbit anti-Bo-LAIT #842 Ab at concentration 2.5 µg/ml in TBST supplemented with 5% skim milk. The blot was rinsed three times for 10 minutes/rinse in TBST. Goat anti-rabbit lgG conjugated with horse radish peroxidase (BioRad) was used to detect rabbit antibody. The membrane was then rinsed three times (10 minutes/rinse) with TBST. The ECL kit (Amersham) was used to visualize the proteins. Lane 1: MW markers; lane 2: nBo-LAIT-842-Sepharose affinity purified nBo-LAIT protein; lane 3: rBo-Sf9- 12CA5 affinity purified, Sf9 insect cell derived recombinant bovine CD14; lane 4: rBo-C127- 842-Sepharose affinity purified, C127 mouse mammary tumor cell line derived recombinant bovine CD14.

Figure 10 shows comparative growth promoting activity of nBo-LAIT and LPS. High buoyant density resting murine splenic B cells were prepared, cultured, and harvested as described for Figure 1A. The indicated concentrations of either affinity purified nBo-LAIT, (affinity purified as described for Figure 1) or LPS, derived from S. typhosa (Difco), were added at the initiation of culture.

Figure 11 shows comparative differentiation promoting activity of nBo-LAIT and LPS. High buoyant density, resting, murine splenic B cells were prepared and cultured as described for Figure 1A. Replicate cultures were initiated using 10 µg/ml of LPS [S.typhosa (Difco)], or 500 ng/ml of affinity purified nBo-LAIT, and harvested at the indicated times. Cumulative lgM production was assessed by quantifying lgM present in supernatants using a commercially available ELISA kit.

Figure 12 shows comparative growth promoting activity of nBo- and rBo-LAIT proteins with that of LPS. High buoyant density, resting, murine splenic B cells were prepared, cultured, and harvested as described for Figure 1A. The indicated concentrations of nBo-LAIT (purified as described for Figure 1A), rBo-LAIT generated in either insect cells, or mammalian cells, and LPS [S.typhosa (Difco)] were added at the initiation of culture. Recombinant Bo-LAIT derived from insect cells was affinity purified from culture supernatants of Sf9 cells transfected with Bo-LAIT cDNA. The expression vector included a 3' 27mer encoding a nonapeptide derived from influenza hemagglutinin (HA tag). Affinity purification

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was achiev d by passing Sf9 supernatants ov r Sepharose conjugated with the mAb 12CA5 (Cell 57:787, 1984), which recognizes the HA tag. Affinity purification of recombinant Bo-LAIT derived from the mainmalian expression system, C127, was achieved as for nBo-LAIT, using Sepharose conjugated with IgG isolated from the polyclonal antiserum derived from rabbit 842.

Figure 13 shows a comparative analysis of the growth promoting activity of nBo-LAIT on purified primary B and T cell populations. High buoyant density, resting, murine splenic B cells were prepared as described for Figure 1A. Primary T cells were isolated from the lymph nodes of the same animals from which the splenic B cells were isolated.

Specifically, lymph node suspensions were passed over anti-Ig columns (Biotex Labora) according to manufacturers instructions, and as previously described (Eur. J. Immunol. 24:967, 1994). T cell populations were >95% CD3+, and <0.5% mIg+, as assessed by immunofluorescent staining and FACS analysis. The left panel shows the proliferative response of cultures containing 1.5x10<sup>5</sup> purified T or B cells in response to the indicated concentrations of affinity purified nBo-LAIT. The right panel shows the proliferative response of the same number of B or T cells to either 50 µg/ml of LPS [S.typhosa (Difco)] or 1 µg/ml of Concanavalin A (Sigma). Serum free conditions were employed, and all stimuli were added at the initiation of culture. At 40 hours, cultures were pulsed with 1 µCi of <sup>3</sup>H-TdR, harvested onto filter discs 6 hours later, and thymidine uptake assessed by scintillation spectroscopy. Numbers indicated represent the average cpm of duplicate cultures.

Figure 14 shows the fetal calf serum independence of nBo-LAIT mediated murine B cell growth. High buoyant density, resting, murine splenic B cells were prepared, cultured, and harvested as described for Figure 1A. Serum free culture medium was supplemented with the indicated concentration of heat inactivated (56°C for 1 hour) fetal calf serum (Gibco BRL) and either no stimulus ( $\blacksquare$ ), 0.5  $\mu$ g/ml affinity nBo-LAIT (O), or 50  $\mu$ g/ml LPS [S.typhosa (Difco)] ( $\blacksquare$ ). At 40 hours, cultures were pulsed with 1  $\mu$ Ci of  $^3$ H-TdR, harvested onto filter discs 6 hours later, and thymidine uptake assessed by scintillation spectroscopy. Numbers indicated represent the average cpm of duplicate cultures.

Figure 15 shows a Northern Blot analysis for CD14 contained in: (1) unfractionated splenocytes, (2) high buoyant density, resting, murine splenic B cells prepared as described for Figure 1A, and (3) FACS sorted mlg <sup>+</sup> B cells derived from the latter high buoyant density resting population. For FACS sorting, high buoyant density cells were incubated with FITC-conjugated mAb 187.1, specific for mouse lgk (Hybridoma 1:5, 1981). The proportion of mlg <sup>+</sup> cells in each of the three populations is indicated. Total RNA was isolated from 10<sup>7</sup> cells of each of the three populations using the Trizol method (Gibco BRL



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Life Technologies) according to the manufacture's instructions and resolved on a 1.2% formaldehyde gel. RNA was transferred to a nylon membrane (GeneScreen) using the vacuum blotting system (Pharmacia). Cross linking, prehybridization and hybridization were performed as recommended by the membrane manufacturer. The murine CD14 specific probe was derived from a genomic mouse CD14 fragment generated by PCR using the forward primer 5'-CTA GAA TTC TCT CCC GCC CCA CCA GAG CCC TGC G-3' (SEQ ID NO:11), and reverse primer 5'-CTA GAA TTC TTA AAC AAA GAG GCG ATC TCC TAG G-3' (SEQ ID NO:12). The amplified fragment was resolved by agarose gel electrophoresis, excised and purified. An L32 cDNA probe (Nucl. Acid Res. 16:10751, 1988), specific for a constitutively expressed mRNA of the large ribosomal subunit protein was used to normalize RNA loading. The probes (100 ng each) were labeled using oligolabeling kit (Pharmacia) to a specific activity of 0.2 - 1x 10° cpm/µg of DNA. The membrane was then washed in 0.2xSSC, 1% SDS at 65 °C for 2 hours and exposed to the X-ray film (Kodak, Biomax MS) for 1 to 5 days.

Figure 16 shows the response of high buoyant density, resting, murine splenic B cells that were further purified for mlg  $^+$  cells as described in Figure 15, or not, to 50  $\mu$ g/ml LPS [S.typhosa (Difco)] and 0.5  $\mu$ g/ml of affinity purified nBo-LAIT. Cultures were established, pulsed with tritiated thymidine, harvested, and thymidine uptake assessed as described for Figure 1A. Numbers indicated represent the average cpm of duplicate cultures.

Figure 17 shows the CD40 independence of recombinant Bo-LAIT mediated B cell activation. High buoyant density, resting, murine splenic B cells were isolated from either conventional C57BL/6 mice, or from CD40 deficient animals created by targeted disruption of the CD40 locus (Immunity 1:167, 1994). Cells  $(1.5 \times 10^5)$  were cultured in the presence of 10  $\mu$ g/ml LPS, or the indicated concentrations of rBo-LAIT derived in the insect cell expression system. Cultures were pulsed and harvested as described in connection with Figure 1A. Numbers indicated represent the average cpm of duplicate cultures.

Figure 18 shows a comparative silver stain (left panel) and immunoblot (right panel) analysis of native CD14 derived from human (nHu), cow (nBo), and mouse (nMo). nHu was isolated from the urine of nephrotic patients, as previously described (Eur. J. Immunol. 24:1779, 1994). nBo was affinity purified from bovine colostrum as described in Figure 3. nMo was isolated from supernatant of the mouse hybridoma OKT3 (PNAS USA 77:4914, 1980) by affinity chromatography using Sepharose 4B immobilized CD14 specific mAb 3C10 (J. Exp. Med. 15:126, 1983). For silver staining, 1 µg of each of the samples was resolved on 10% SDS-PAGE at 200V for 45 minutes. Protein was visualized by silver staining (Biorad) following manufacturers instructions. For immunoblotting, 250 ng of each of the samples was resolved by 10% SDS-PAGE at 180 mA for 45 minutes, electrophoretically transferred to

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PVDF m mbrane (Millipor ) at 180 mA for 30 minutes. The m mbrane was blocked for 1 hour in 5% skim milk in TBST (20mM Tris.HCl, pH 7.5, 150mM NaCl, 0.025% Tween 20), followed by incubation for 1 hour with mAb 3C10 (J. Exp. Med. 15:126, 1983) at concentration 10 µg/ml in TBST supplemented with 5% skim milk. The blot was rinsed three times for 10 minutes/rinse in TBST. Goat anti-mouse IgG conjugated with horse radish peroxidase (BioRad) was used to detect mouse antibody. The membrane was then rinsed three times (10 minutes/rinse) with TBST. The ECL kit (Amersham) was used to visualize the proteins.

Figure 19 shows a comparative analysis of native CD14 derived from human (nHu), cow (nBo), and mouse (nMo) to stimulate the growth of murine B cells. These three proteins were purified as described in Figure 18. The response of high buoyant density, resting, murine splenic B cells isolated as described in Figure 1A, to the indicated concentrations of the three proteins, was assessed. Cultures were pulsed with 1µCi of <sup>3</sup>H-Tdr at 40 hrs and harvested at 46 hrs. The numbers indicated represent the index of stimulation derived by dividing mean cpm of duplicate cultures stimulated with the indicated concentration of nHu, nBo, and nMo, by mean cpm of duplicate cultures containing no stimulus.

Figures 20 A and B shows growth promoting activity of nBo-LAIT on human B cells isolated from cord blood and tonsils, respectively. Figure 20A shows thymidine uptake by cord blood B cells isolated by positive selection. Cord blood leukocyte suspensions were stained with fluorescein labeled mAb specific for the pan B cell marker CD72. CD72 positive cord leukocytes were then isolated on a fluorescence activated cell sorter (FACStar Plus, Becton Dickenson) resulting in purities of >98%. B cells (1.5x10<sup>5</sup>) were cultured as described for Figure 1A, in the presence of no stimulus or 2  $\mu$ g/ml of nBo-LAIT. B cells were also cultured in wells which had been pre-coated for 9 hours with a combination of two mAbs, one specific for human IgK (LO-HK3, (In "Rat Hybridomas and Rat Monoclonal Antibodies" ed. H. Bazin, CRC Press, Boca Raton, FL, USA)] and one specific for human Igλ [LO-HL2, (In "Rat Hybridomas and Rat Monoclonal Antibodies" ed. H. Bazin, CRC Press, Boca Raton, FL, USA)], each at a coating concentration of 1.5  $\mu$ g/ml, without additional stimulus, or in the presence of 2µg/ml of nBo-LAIT. Cultures were pulsed at 60 hours with 1 µCi of 3H-TdR, harvested onto filter discs 12 hours later, and thymidine uptake assessed by scintillation spectroscopy. Figure 20B shows results obtained using tonsil B cells prepared by negative selection. Specifically, leukocyte suspensions were labeled with biotinylated mAb specific for CD3s (Becton Dickenson), followed by avidin conjugated with iron containing "micro-beads" (Becton Dickenson). The labeled population was passed through the MACS (Becton Dickenson), and the effluent collected. This population contained <1% T cells, and >97% B cells as assessed

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by immunofluorescence staining with lineage specific mAbs. B cells (1.5 x  $10^5$ ) were cultured as described in connection with Figure 1A. As for cord blood B cells, tonsil B cells were cultured in the presence and absence of plate bound mAbs specific for human lgK and  $\lambda$ , but in this case, wells were pre-coated using a concentration of 0.5  $\mu$ g/ml of each of the mAbs. Cultures were pulsed, harvested, and thymidine uptake assessed as described for Figure 20A.

Figure 21 shows the concentration of CD14 in breast milk over time post-partum. Breast milk was collected from two donors (A.D. and S.B.) at the indicated times post-partum. Clarified whey was prepared, and the concentration of contained CD14 assessed using a CD14 specific ELISA kit (IBL, Hamburg) according to manufacturers instructions.

#### 10 DESCRIPTION OF PREFERRED EMBODIMENTS

The experiments described below demonstrate purification of native bovine LAIT protein (nBo-LAIT), also referred to herein as bovine CD14, from colostral whey. Amino acid sequence analysis of purified nBo-LAIT is shown, and homology with human CD14 is demonstrated. A method for the purification of human CD14 from is shown. A method for the purification of mouse CD14 from hybridoma supernatant is shown.

In vitro B cell stimulation assays are described for affinity purified colostral Bo-LAIT, human colostral CD14, human CD14 derived from urine, and mouse CD14 derived from a hybridoma supernatant. High buoyant density resting splenic B cells derived from mouse are shown to enter and progress through cell cycle, in response to LAIT protein from the three species, and to differentiate into high rate immunoglobulin secreting cells in response to exposure to LAIT-protein from bovine. These activation events occur in defined serum free medium, and it is also shown that the presence of fetal calf serum in culture medium does not affect LAIT protein mediated B cell activation. Experiments are shown demonstrating that nBo-LAIT specifically activates murine B cells, and not murine T cells, and that LAIT protein activates B cell populations in which CD14 mRNA is undetectable. Experiments demonstrating that bovine CD14 induces growth of B cells in which CD40 is not expressed are also given.

The isolation, cloning and sequencing of both genomic DNA and cDNA encoding bovine CD14 is described. Sequence comparisons with mouse and human CD14s, known in the literature, show the sequence relationship between Bo-LAIT and these previously known CD14s. B cell growth and differentiation activities associated with recombinant bovine CD14 are shown.

Methods for the expression of recombinant bovine CD14 in both insect and mammalian systems are described. Specifically, a baculovirus expression vector was employed in aid of expressing recombinant proteins in insect cells. Comparison of the B cell

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growth and differentiation properties of native Bo-LAIT (nBo-LAIT) and recombinant Bo-CD14 (rBo-LAIT) derived from the baculovirus expression system revealed that the latter was functional, and had a specific activity of roughly 50% of that of nBo-LAIT.

The mouse mammary carcinoma cell line, C127, was used as a recipient of cDNA encoding CD14 derived from bovine. cDNA was cloned into a bovine papilloma virus expression vector. Stable C127 transfectants were established, and recombinant CD14 protein was isolated from supernatants of confluent C127 cultures by affinity chromatography. Western blot analyses of insect cell and C127 derived recombinant LAIT-proteins revealed that different co- and/or post-translational modifications were generated in the two expression systems. The specific activity of mammalian cell derived recombinant bovine CD14 was the same as that recombinant material derived from insect cells.

A comparison of the B cell growth promoting activity supported by native Bo-LAIT and recombinant bovine CD14 derived from insect cells and mammalian cells is given. Further, growth promoting activity of native Bo-LAIT activities on human B cells, isolated from either tonsils, or from cord blood, is given. Results demonstrate that as for murine B cells, human B cells, isolated from either from a neonate or an adult, are susceptible to Bo-LAIT mediated growth.

It is shown that the concentration of CD14 present in human colostrum, and in breast milk up to 78 days post-partum, is between 3-20-fold higher than that observed in sera from healthy donors.

#### **METHODS**

### Purification of Bovine LAIT-protein

More than five liters of colostrum was obtained from the first mammary secretions of cows having just given birth.

(i) Clarified colostral whey was prepared by centrifugation of colostrum first at 4420g for 30 minutes to remove cells and cellular debris. The supernatant of this spin was then centrifuged at 250,000g for two hours. The floating lipids and the pelleted casein were discarded, and the clarified colostral whey was subjected to further fractionation.

Each fraction derived from each fractionation technique was assessed for B cell growth promoting activity in vitro. Thus, each fraction was assayed over a wide concentration range for its capacity to stimulate the growth of high buoyant density, resting B cells derived from mouse spleen, as previously described (J. Immunol. 131:581, 1983). Defined serum free medium was used throughout these analyses [IMDM (Gibco), supplemented with 5 x  $10^{-5}$  2- $\beta$ -mercaptoethanol, 5  $\mu$ g/ml iron-saturated transferrin

(Boehringer, L w s, GB), 0.5 mg/ml delipidated BSA (Boehringer), 100 U/ml penicillin (Gibco), 100  $\mu$ g/ml streptomycin (Gibco), and essential amino acids). Fractions derived from the isolation scheme described below were tested directly, as well as in combination with a submitogenic concentration of LPS (0.25  $\mu$ g/ml). As will be described, as LAIT protein approached purity, its direct mitogenic properties were revealed.

(ii) Salting out of proteins contained within colostral whey preparations was accomplished using sequential precipitation in (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>. The sequence of increasing salt concentrations employed was 42%: 50%: 62%: 65% (v/v) ammonium sulphate (AS). Thus, the concentration of AS in the supernatant of the material precipitated at 42% was increased to 50%; the material precipitated at 50% rescued, and the concentration of AS in the remaining supernatant increased to 62%, and so on. Each AS precipitated pellet was solubilized in 10mM Tris-HCL pH 8.0, containing 0.15M NaCl and 1 mM AEBSF (TNAEBSF). These fractions were desalted and buffer exchanged to TNAEBSF using 10DG columns, and assayed for bioactivity. The majority of B cell growth promoting activity was isolated in the 62% AS precipitate following the above scheme (not shown).

(iii) Activity was subsequently enriched, and ultimately purified using three sequential protein fractionation techniques. Fifty milligrams of the 62% AS enriched fraction was applied to an anion exchange column, and the material separated using a salt gradient of 50mM to 400mM NaCl in 10mM Bis-tris propane, with a simultaneous pH gradient of 7.5 to 9.5. Figure 1A shows the elution profile from this column, and Table 1A indicates the fraction containing the peak activity, fraction #57. Twenty milligrams of fraction #57, was then applied to a molecular sieving column equilibrated in 20mM Tris-HCL, pH 8.0 containing 0.45M NaCl. The elution profile of this fractionation is shown in Figure 1B and the activity of the peak fraction #38 shown in Table 1B.

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TABLE 1A	CPM x 10 <sup>-3</sup>
NO STIMULUS	0.8
LPS 50 µg/ml	152.5
LPS 0.25 µg/ml	3.9
FRACTION 57 + 0.25 µg/ml LPS	108.7

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TABLE 1B	CPM x 10 <sup>-3</sup>
NO STIMULUS	0.4

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LPS 50 µg/ml	102.1
LPS 0.25 µg/ml	1.3
FRACTION 36 + 0.25 µg/ml LPS	76.0

TABLE 1C	CPM x 10 <sup>-3</sup>
NO STIMULUS	0.7
LPS 50 µg/ml	135.2
LPS 0.25 µg/ml	3.5
FRACTION 25 + 0.25 µg/ml LPS	112.0

One milligram of fraction #38 was then applied to an hydroxy apatite column in 1mM NaCl, and eluted using a gradient of 1 to 500mM potassium phosphate buffer pH 6.8. The elution profile is shown in Figure 1C with the associated activity shown in Table 1C.

The inset in Figure 1C represents an SDS-PAGE analysis of the fraction with peak activity followed by silver staining, and illustrates a single major band with a relative molecular mass of 46-50kD.

## 15 Sequence Analysis of Bovine LAIT-protein

The purified nBo-LAIT was subjected to sequence analysis. The N-terminus was found to be blocked. The material was subjected to hydrolysis with either cyanogen bromide, or trypsin. Five fragments were generated and these were purified using either reverse phase HPLC, or SDS-PAGE followed by electroblotting onto a PVDF membrane, prior to sequencing.

As illustrated in Figure 2, the five fragments all aligned, with significant homology, to human CD14.

### Affinity Purification of LAIT Protein from Bovine and Human Colostrum

nBo-LAIT isolated using classical protein fractionation techniques was used to prepare a rabbit (#842) polyclonal antibody. The IgG fraction of this antiserum was purified on Protein A-Sepharose, and subsequently conjugated to Sepharose 4B.

The sequence homology of nBo-LAIT and human CD14 (HuCD14) indicated that Bo-LAIT might be the bovine homologue of CD14. This was further explored by generating an affinity column using available monoclonal antibody (mAb) specific for HuCD14. This antibody, 63D3 (PNAS 77:6764, 1980), was purified from the corresponding hybridoma supernatant on an affinity column comprised of mAb 187.1 [ rat anti-mouse kappa (Hybridoma 1:5, 1981)], conjugated to Sepharose 4B, and the purified mAb then conjugated to Sepharose

4B.

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Bovine clarified colostral whey was sequentially salted out using ammonium sulphate, as described above. Human colostral whey was fractionated on a Sephacryl S100 HR column. The fractions containing peak B cell growth promoting activity were then affinity purified using either the 842-Sepharose column for the bovine material, or the 63D3-Sepharose column for the human material.

The SDS-PAGE analysis of affinity purified colostral Bo-LAIT, and affinity purified human colostral CD14 is shown in Figure 3 and the associated B cell growth promoting activity is shown in Table 2. As illustrated, a predominant band was isolated from both colostral preparations, the p46-50 bovine material (Figure 3, lane 3) and a p50-52 human molecule (Figure 3, lane 4).

TABLE 2	BOVINE #842 pH2.5 ELUATE	HUMAN 6303 pH2.5 ELUATE
100 ng/ml	20.00*	0.3
10 ng/ml	1.49	0.3
• + LPS1	16.5	21.1
• + b-7-6	5.17	8.5

\* Numbers represent cpm x 10<sup>-3</sup> at 40 hours of culture.

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The bioactivity shown in Table 2 demonstrates that affinity purified Bo-LAIT, at a concentration of 100 ng/ml, stimulated the growth of resting mouse B cells. When added at 10 ng/ml, this material was no longer mitogenic, but costimulation was achieved upon the addition of either a submitogenic concentration of LPS, or a mAb specific for mouse IgM, b-7-6 (Eur. J. Immunol. 14:753, 1984). The affinity purified human material was not by itself found to be mitogenic at concentrations tested, but at 10ng/ml, B cell growth was stimulated with the same costimuli as efficiently as with the bovine material.

The bioactivity of nBo-LAIT is heat labile. As illustrated in Figure 4A, treatment of affinity purified Bo-LAIT at 95°C for 10 minutes abolishes the associated B cell growth promoting activity. Similar treatment of LPS had no effect on its activity (inset of Figure 4A). Further, the polyclonal anti-Bo-LAIT, #842, efficiently blocked the B cell growth promoting activity of nBo-LAIT, while not affecting the activity of LPS. See Figure 4B and inset.

### Molecular Cloning of Genomic Bovine CD14

A bovine genomic EMBL3 SP6/T7 lambda library (Clontech) was screened with

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a 1.5 kb fragment of human CD14 cDNA (obtained from R.Ulevitch, Scripps Institute). Fifteen positive signals were obtained, and the strongest signal, clone "B2" was chosen for further analysis and cloning of bovine CD14.

Isolated and purified phage DNA from clone B2 had an insert size of roughly 15 kb. Purified DNA was digested, and a resulting 7.1 kb EcoRI-Xhol fragment, containing a homologous sequence to human CD14, was subcloned into pBluescriptSK + (Stratagene). Restriction mapping, utilizing a wide range of enzymes, followed by hybridization with the human CD14 probe enabled the localization of the bovine CD14 gene within the cloned fragment (Figure 5A). Further restriction mapping was used for the subcloning of four shorter fragments (I-IV) into pBluescriptSK +, and the subsequent sequencing of roughly 5 kb encompassing the entire bovine CD14 gene (Figure 5C). Fragment I (EcoRI-BamHI, 3.2kb); Il (PstI-PstI, 1.35kb); Ill (PstI-PstI, 0.3kb); and IV (PstI-XhoI, 0.95kb), were used to construct nested overlapping unidirectional deletions. These fragments provided contiguous sequence of the bovine CD14 locus. Figure 5B depicts the organization of the bovine CD14 genomic fragment.

### Molecular Cloning of Bovine CD14 cDNA

Poly(A +) RNA was isolated from bovine peripheral blood monocytes, and Gigapack II Packaging Extract (Stratagene) was used to package recombinant lambda phage DNAs. A cDNA library was prepared using the Excell EcoRI/CIP vector with the "Time Saver cDNA Synthesis Kit" (Pharmacia).

The library was screened with the probe derived from the coding translated region of the bovine genomic CD14 fragment by PCR (details are provided below in the section describing the preparation of baculovirus recombinant expression vector with bovine CD14 fragment). The probe was labeled with <sup>32</sup>P by random hexanucleotide-primed second strand synthesis (Oligolabelling Kit, Pharmacia Biotech). Screening procedures were performed under conditions of high stringency (0.1xSSC, 1%SDS, 65°C for 3 hours). One of the clones obtained (ExCell/BoCD14-1), contained a 1.4kb insert, which was subcloned into pBluescript SK +, and sequenced using pBS/BoCD14 subclones containing progressive overlapping unidirectional deletions (Nested Deletion Kit, Pharmacia).

This bovine CD14 cDNA clone consists of 1327bp. An ATG initiation codon is followed by an open reading frame of 1116 nt, and a TAA stop codon at nucleotide 1202. The open reading frame is flanked by 82bp of 5' untranslated sequence and 122bp of 3' untranslated sequence. A polyadenylation signal, 5'-ATTAAAA-3', is located 105bp 3'of the termination codon.

Alignment of bovine CD14 genomic and cDNA sequences reveals that they are colinear from the start of 5' cDNA until the first and only intron (88bp) which is found immediately after the ATG initiation codon. The remainder of the coding sequence is

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uninterrupted. Thus, the intron-exon organization pr viously described for human and mouse CD14 is precisely conserved in bovine CD14. Comparison of the translated nucleotide sequence of bovine CD14 cDNA with those of human and mouse CD14 cDNAs revealed 74.2% and 62.6% nucleotide identity in coding regions, respectively (Figure 6).

The primary structure of the bovine CD14 protein was deduced from cDNA sequence, and consists of 374 amino acids. The first methionine is followed by a stretch of 15 hydrophobic and/or neutral residues, typical of eukaryotic signal peptides. Alignment of the amino acid sequences of bovine CD14 with human and mouse CD14 reveals 73.1% and 62.3% identity, respectively (Figure 7). There are three potential N-linked glycosylation sites (Asn-X-Thr/Ser) all of which are conserved in human and mouse CD14. Moreover, bovine CD14 contains 10 leucine-rich repeating motifs (LXXLXL), common to both human and mouse CD14 (J. Immunol. 145:331, 1990).

### **Expression of Recombinant Bovine CD14 in Insect Cells**

In preparing DNA fragments for producing recombinant CD14 proteins, full length fragments of CD14 translated regions were generated by PCR. Specific sets of PCR primers were designed based on sequence information obtained from bovine CD14 cDNA. The PCR primer for the 5' end contained: two recognition sequences for Nhel; a Kozak sequence; an ATG initiation codon; and the first 17-21 nucleotides of translated coding region. The PCR primer for the 3' end contained: two recognition sequences for Nhel; and the last 21-24 nucleotides of translated coding region up to and excluding the TAA stop codon (Figure 8A).

The bovine CD14 translated region was amplified using the 7.1kb EcoRl-Xhol genomic CD14 fragment (see above) as a template. PCR was carried out using Pwo DNA polymerase (Boehringer). Amplification was done by adding 5ng of template DNA , 10 mM Tris-HCl pH 8.85, 25 mM KCl, 5mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 2mM MgSO<sub>4</sub>, 250 mM of each dNTP, 250 nM of each primer, and 5 units of Pwo DNA polymerase, in a final volume of 100  $\mu$ l. The samples were amplified for 30 cycles at 70°C annealing temperature using a DNA Thermal Cycler (Perkin Elmer).

Amplified fragments were digested with Nhel, and individually subcloned downstream of the polyhedrin promoter in the baculovirus transfer vector pETL-HA (C. Richardson, OCI/Amgen). This vector is derived from pETL (Methods in Molecular Biology 39:161, 1995), and contains a 3' 30bp Nhel-BamHI DNA fragment encoding a nonapeptide derived from influenza hemagglutinin (HA), followed by the stop codon TAG (5'-TAC CAA TAC GAT GTT CCA GAT TAC GCT TAG-3')(SEQ ID NO:13). The recombinant transfer vectors were individually cotransfected with the wild-type baculovirus Autographa californica nuclear polyhedrosis virus (AcMNPV, Linear Transfection Module, Invitrogen) into Sf9 cells (Methods in Molecular Biology 39:161, 1995). The recombinant baculovirus clones were selected and purified according to established protocols (Methods in Molecular Biology 39:161, 1995). Sf9

cells were infected with recombinant baculovirus at a multiplicity of 5-10:1.

A time course analysis was performed to determine the optimum time period required for the infected Sf9 cells to secrete recombinant CD14 proteins. Immunoblot analysis of the cell media taken at different time points using the anti-HA monoclonal antibody 12CA5 (Cell 57:787, 1984), revealed that the expression of recombinant CD14 proteins reached the maximum level at 96 hours. This period was chosen in subsequent experiments for the production of recombinant proteins for bioassay (see below). Western blot analysis of Sf9 derived recombinant bovine CD14 is illustrated in Figure 9.

### Expression of Recombinant Bovine CD14 in Mammalian Cells

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We used a modified version of the pBPV Episomal Mammalian Expression Vector (Pharmacia) for stable expression of recombinant bovine CD14 in mammalian cells. To enable direct selection of transformed cells, pBPV was modified by including a neomycin resistance gene, which was inserted 3.4kb upstream of the expression cassette. Towards this end, a 1.95kb HindIII-Xbal fragment from pBCMGSneo (Eur. J. Immunol. 18:98, 1988) was subcloned into pCRII (Invitrogen). The recombinant construct, pCRII-neo, was purified, and the cloned fragment was amplified by PCR. PCR primers were designed such that the recognition sequence for Sal I was included at both the 5' and 3' ends. Primer sequences were complementary to the polylinker region of pCRII vector, flanking the HindIII (Primer A) and Xbal (Primer B) cloning sites.

Primer A: 5'-GCA GTC GAC ACT ATA GAA TAC TCA AGC-3'
(SEQ ID NO:14)

Primer B: 5'-TTC GTC GAC ATT GGG CCC TCT AGA-3' (SEQ ID NO:15)

The final product was digested with Sal I, gel purified, and subcloned into the Sal I cloning site of pBPV in the same transcription orientation as that of the contained expression cassette. Plasmid preparations of the modified expression vector, pBPVneo-13, were generated (Plasmid Maxi Kit, Quiagen).

A DNA fragment encoding the translated region of bovine CD14 was prepared by PCR amplification of the gene in the pETL-HA vector. The 5' end PCR primer used in the amplification reaction included: an Xho I recognition sequence; followed by an Nhe I recognition sequence (present in the pETL-HA vector); a Kozak sequence; an ATG initiation codon; and the first 11 to 13 nucleotides of the translated region. The core PCR primer for the 3' end contained the HA coding sequence which was extended, as for the 5' sequences, with the inclusion of an Xho I recognition sequence. Primers are shown in Figure 8B.

PCR was carried out using Pwo DNA polymerase (Boehringer Mannheim).

Amplifications were done by adding 5ng of DNA template, 10 mM Tris-HCl pH 8.85, 25 mM

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KCI, 5mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 2mM MgSO<sub>4</sub>, 250 mM of each dNTP, 250 nM of each primer, and 5 units of Pwo DNA polymerase in a final volume of 100  $\mu$ l. The samples were amplified for 30 cycles at 70°C annealing temperature using a DNA Thermal Cycler (Perkin Elmer).

The amplified fragments were digested with Xho I and gel purified. These fragments were then subcloned into pBPVneo-13 downstream of the mouse metallothionein I promoter. Prior to the subcloning, pBPVneo-13 was pretreated with the appropriate restriction enzyme, and dephosphorylated using calf intestinal phosphatase. Recombinant plasmids were prepared using a Plasmid Maxi Kit (Quiagen).

The recombinant plasmid (pBPVneo13-BoCD14) was transfected into the mouse mammary tumor cell line, C127 (PNAS 78:2727, 1981), using 20  $\mu$ g of DNA/10 $^{7}$  cells. DNA transfer was achieved by electroporation at 960  $\mu$ F/280V using a Gene pulser (Bio-Rad Laboratories). Stable transformants were selected in the presence of 1.5 mg/ml G418 (Life Technologies).

Transfectants expressing high levels of membrane CD14 (non transfected C127 are negative for CD14) were enriched by immunofluorescence staining followed by fluorescence activated cell sorting using a Becton Dickinson FACStar Plus. The level of membrane expression of the exogenous protein correlated well with the amount of secreted CD14 rescued in 48 hour supernatants of confluent cultures of transfected C127 cells. Unlike the purification of recombinant material generated in insect cells, it was not found possible to affinity purify C127 derived material using 12CA5-Sepharose affinity columns. This might have been due to the loss of the C-terminal HA tag on recombinant proteins derived from C127 cells. As a consequence, recombinant bovine CD14 derived from C127 was affinity purified on 842-Sepharose. Immunoblot analysis of recombinant bovine CD14 derived from C127 cells is illustrated in Figure 9.

### Comparative Growth and Differentiation Promoting Activities of nBo-LAIT and LPS

The results shown in Figure 10 illustrate that native Bo-LAIT supports the growth of high buoyant density, resting, murine splenic B cells with efficiencies roughly 200-fold higher than that of LPS. Thus, nBo-LAIT at 50 ng/ml results in the induction of DNA synthesis comparable to that observed in the presence of 10  $\mu$ g/ml of LPS.

The capacity of nBo-LAIT to induce B cell growth is paralleled by its capacity to induce the differentiation of high buoyant density, resting, murine B cells to immunoglobulin secretion. As illustrated in Figure 11, the amount of cumulative IgM induced by 500 ng/ml of nBo-LAIT is comparable to that induced by 10  $\mu$ g/ml of LPS. The amount of IgM secretion within a 24 hour culture period was assessed. 500 ng/ml of nBo-LAIT induced 956±10 ng/ml; 754±8.7 ng/ml; and 25±1.4 ng/ml of IgM within the 24 hour culture periods of 48-72 hours; 72-96 hours; and 96-120 hours, respectively. Corresponding values derived from cultures stimulated with 10  $\mu$ g/ml of LPS were: 1442±71 ng/ml; 874±32 ng/ml; and 183±3 ng/ml, respectively. Thus, nBo-LAIT has the capacity to induce high buoyant density,

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resting murine B cells to immunoglobulin secretion at rates comparable to those observed when the B cells are stimulated with the most potent stimulus currently known. Further, it has the capacity to do so at concentrations of 1-10% of that of LPS.

The capacity of nBo-LAIT to induce isotype switching of murine resting B cells was also assessed. The supernatants derived from the cultures described above were also assessed for the presence of IgG isotypes. It was observed that 500 ng/ml of nBo-LAIT and  $10 \,\mu\text{g/ml}$  of LPS induced cumulative levels (ng/ml at day 5) of IgG1:  $7.0 \pm 0.1$  and  $5.6 \pm 0.6$ ; IgG2a:  $358 \pm 3$  and  $406 \pm 8$ ; IgG2b:  $8 \pm 1$  and  $11 \pm 2$ ; IgG3:  $75 \pm 5$  and  $75 \pm 0.5$ ; and IgA:  $6.5 \pm 1.5$  and  $5.0 \pm 0.3$ , respectively. Thus, nBo-LAIT, has the capacity, to induce some isotype switching by resting murine B cells in the absence of T cells.

### Comparative Growth Promoting Activities of nBo- and rBo-LAIT Proteins

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Recombinant forms of bovine CD14, both those derived in insect cells and mammalian cells, have the capacity to induce the growth of high buoyant density, resting, murine B cells. As illustrated in Figure 12, rBoCD14 derived from insect cells, and affinity purified on 12CA5-Sepharose, induces robust DNA synthesis at 0.2-3 µg/ml concentrations. Comparible activity was supported by recombinant material derived from the mammalian expression system, and both recombinant forms support B cell growth at ng/ml concentrations, comparable to the activity observed for colostrum derived nBo-LAIT.

Assurance that the bioactivity mediated by nBo-LAIT isolated from bovine colostrum either by classical protein fractionation techniques, or affinity purification, is mediated by the observed protein, comes from the assessment of the bioactivity mediated by recombinant bovine CD14. As illustrated in Figure 9, the apparent molecular weights of neither of the recombinant forms of bovine CD14 are identical to that of nBo-LAIT. The reason for the observed differences in apparent molecular weight is not clear but might be due to either distinct co- and/or post-translational modifications, distinct sizes of the core proteins, or both. Monocyte derived soluble CD14 has been documented and can be generated in one of three currently understood mechanisms, each of which would result in proteins of distinct molecular weight. It can be secreted as a full length molecule (Eur. J. Immunol. 23:2144, 1993; Eur. J. Immunol. 25:604, 1995), the membrane expressed GPI linked form can be cleaved by phospholipases (J. Immunol. 141:547, 1988; EMBO J. 13:1741, 1994), and the membrane expressed GPI linked form can be cleaved by serine/threonine proteases, putatively expressed on the outer plasma membrane of the monocyte it self, and activated in as yet uncharacterized ways (J. Immunol. 147:1567, 1991). It remains to be determined whether the distinct apparent molecular weights of rBo-LAITs and colostrum derived nBo-LAIT is due to distinct co- and/or post-translational modification of the recombinant materials supported by their respective expression systems, or distinct core protein sizes, or both.

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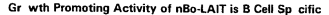
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Having observed the bioactivities of nBo-LAIT on murine B cells, effects on the physiology of murine T cells was examined. The fact that isolated purified B cells do respond to nBo-LAIT does not preclude the possibility that Bo-LAIT can also activate T cells.

Lymph node T cells were isolated by negative selection on anti-Ig columns (Biotex Labora) as previously described (Eur. J. Immunol. 24:967,1994). The resuting effluent population contained >95% CD3+ cells and >0.5% mlg+ cells as assessed by immunofluoresence staining and FACS analysis. High buoyant density splenic B cells were isolated from the same mice, and both the lymph node T cells and splenic B cells assessed for their responsiveness to nBo-LAIT. The purity of these populations was assessed functionally by analyzing their responses to the B cell specific mitogen LPS, and the T cell specific mitogen, concanavalin A (ConA). As illustrated in the right panel of Figure 13, the B cells, but not the T cells responded with robust DNA synthesis to LPS, while the T cells, but not the B cells responded to ConA. As illustrated in the left panel, the T cells did not respond to nBo-LAIT over the dose range tested, while the B cells responded to nBo-LAIT at concentrations of 10 ng/ml and higher.

#### LAIT-protein Induced B Cell Growth and Fetal Bovine Serum

It has been demonstrated recently that monocytes respond to soluble CD14 (sCD14) isolated from the urine of nephrotic patients with the production of pro-inflammatory cytokines (Eur. J. Immunol. 24:17790,1994), in the absence of serum derived lipopolysaccharide binding protein (LBP). In contrast, the capacity of low concentrations of LPS to stimulate the production of these same cytokines by CD14 expressing monocytes is serum dependent (J. Exp. Med. 176:719, 1992). This serum dependence is overcome at high concentrations of LPS. Thus, cytokine production by monocytes induced by 10 ng/ml LPS is strictly dependent on serum/LBP, but that induced by 50  $\mu$ g/ml LPS is not.

To determine whether the presence of fetal bovine serum LBP would affect the capacity of low concentrations of nBo-LAIT to induce the growth of high buoyant density murine B cells, the response elicited by 500 ng/ml nBo-LAIT was assessed over a broad range of FBS concentrations, in comparison to that induced by 50  $\mu$ g/ml LPS. As illustrated in Figure 14, the B cell responses to both these stimuli were unaffected by the presence of up to 9% FBS present in the culture medium.

# LAIT-protein Induced B Cell Growth and mCD14

The diffential sensitivity of B cells and monocytes to LPS mediated activation could be related to the expression of membrane CD14 (mCD14) by the latter. It has been directly demonstrated that the sensitivity of LPS mediated activation can be dramatically enhanced by the presence of mCD14 on responding cells. Specifically, it has been

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demonstrated that the sensitivity of a mCD14 negative pre-B cell line to LPS is increased by roughly a thousand-fold upon the forced expression of exogenous GPI linked mCD14 (J. Exp. Med. 175:1697,1992). Whether primary B cells express mCD14 remains contentious. The high specific activity of Bo-LAIT relative to that of LPS in B cell activation is consistent with its mode of activation being independent of mCD14 expression.

To directly assess the involvement of mCD14 expression in Bo-LAIT mediated B cell activation, the presence of CD14 specific mRNA was assessed in the high bouyant density murine B cell populations used as responders. Total RNA from 10<sup>7</sup> cells from each of three spleen derived populations: unfractionated splenocytes; T-depleted, Percoll fractionated, high buoyant density cells, those used routinely in LAIT-protein mediated growth assays; and membrane Ig expressing cells isolated from the high buoyant density T-depleted population by immunofluorescence labelling with FITC conjugated rat mAb specific for mouse Igk [187.1 (Hybridoma 1:5, 1981)) and subsequent purification using a Becton Dickinson FACStar Plus cell sorter. As illustrated in Figure 15, these three populations contained 59.2%; 83.5%; and 99.8% mlg + cells. Isolated RNA from these populations was resolved on a 1.2% formaldehyde gel, and transferred to a nylon membrane (GeneScreen), crosslinked, prehybridized, and hybridized with two probes, sequentially.

The mouse CD14 probe was derived by PCR from genomic DNA. Specifically, amplification was done by adding Pwo DNA polymerase (Boehringer Mannheim) to 0.4  $\mu$ g of Balb/c genomic DNA with the forward primer: 5'-CTA GAA TTC TCT CCC GCC CCA CCA GAG CCC TGC G-3' (SEQ ID NO: 11); and reverse primer: 5'-CTA GAA TTC TTA AAC AAA GAG GCG ATC TCC TAG G-3' (SEQ ID NO: 12). The sample was amplified for 30 cycles in a DNA thermal cycler (Perkin Elmer) using an annealing temperature of 55°C. The amplified fragment was resolved by agarose gel electrophoresis, excised from the gel, and purified. An L32 probe (Nucl. Acid. Res. 16: 10751, 1988) was used to normalize RNA loading. Each probe (100 ng) was labelled using an oligolabelling kit (Pharmacia) to a specific activity of 0.2-1x10° cpm/ $\mu$ g DNA. Membranes were probed and washed in 0.2xSSC, 1% SDS at 65°C for 2 hours, and exposed.

As illustrated in Figure 15, RNA derived from both the unfractionated splenocytes, and the T-depleted high buoyant density splenocytes contained CD14 specific mRNA, while the FACS purified 8 cells did not. The CD14 signal in the T-depleted, high buoyant density splenocytes, heretofore referred to as resting 8 cells, is likely due to contaminating monocytes, as these populations are 85-90% mlg +, but > 98% MHC Class II + as assessed by immunofluoresence staining and FACS analysis. It is therefore possible that 8o-LAIT mediated growth of 8 cells in this population is indirect, and is mediated through the activation of contained monocytes.

The response of T-depleted, Percoll fractionated splenocytes, and FACS purified mlg <sup>+</sup> cells derived from this population to nBo-LAIT, and LPS induced growth was compared.

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As illustrated in Figure 16, both populations responded robustly to both stimuli. The 10-fold higher stimulation indices obtained with 99.8% pure B cell populations was due to the reduced non-stimulated background observed in these cultures (Figure 16).

### LAIT-protein Induced B Cell Growth and CD40

As described above, mAbs specific for CD40, which is expressed on the membrane of B cells, have been observed to induce growth of murine B cells (Sem. in Immunol. 6:267, 1994; PNAS 83:4494, 1986; J. Immunol. 140:1425, 1988;).

To determine whether there is some relationship between anti-CD40 and LAIT-protein induced B cell activation, the capacity of rBo-LAIT to stimulate the growth of high buoyant density splenic B cells isolated from either conventional C578L/6 mice or C578L/6 mice in which CD40 expression was ablated through targeted gene disruption was examined (Immunity 1:167, 1994). As illustrated in Figure 17, no differences in the responses of these B cells were observed over the concentration range of rBo-LAIT tested.

These results indicate that CD40 per se need not be involved in LAIT-protein signalling, but the possibility that second messenger generating systems utilized by CD40 and the putative membrane receptor for LAIT-protein are shared, cannot be excluded.

# Comparative Analysis of native Human, Bovine and Murine CD14

Having observed the activities of nBo-LAIT and rBoCD14 on murine B cells, the activities of mouse CD14 (Mo) and human CD14-(Hu) isolated from fluids other than colostrum were examined. It has been demonstrated that Hu-CD14 is present in the urine of nephrotic patients (Eur. J. Immunol. 24:1779,1994). Hu-CD14 was isolated using a modified protocol. Urine was precipitated by adding saturated (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> to a final concentration of 45% (v/v), and precipated material cleared by centrifugation at 14000g for 30 minutes. (NH<sub>a</sub>)<sub>2</sub>SO<sub>4</sub> concentration in the supernatant of this spin was then increased to 75% (v/v). The precipitate was pelleted at 14000g for 30 minutes, and solubilized in TN buffer pH8.0 containing 10mM Tris, 150mM NaCl, and "Complete" protease inhibitor cocktail (Boehringer Mannheim). Insoluble material was cleared by centrifugation at 13000g for 15 minutes. The supernatant of this spin was desalted on G-10 columns (Bio Rad) equilibriated in TN buffer pH8.0. This material was then passed over Sepharose 4B to which the human CD14 specific mAb, 3C10 (J. Exp. Med. 15:126,1983), had been conjugated. Bound material was eluted in 100mM acetate, 150mM NaCl, pH2.8, and immediately neutralized by adding a tenth volume of 1M Tris, pH 8.0. The eluate was concentrated in a speed-vac, and protein concentration determined colorimetrically.

Murine CD14 was isolated from the supernatant of the mouse hybridoma OKT3 (PNAS USA 77:4914,1980). During a screening analysis of cell populations for the expression of CD14 specific mRNA, it was observed that every hybridoma assayed contained message. It

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follows that if the donor and fusion partner were both of murine origin, the CD14 produced would also be of murine origin. The hybridoma OKT3 satisfies these criteria. To assess whether CD14 protein was being produced by OKT3, and in sufficient quantities to allow isolation, material contained in 1 litre of OKT3 culture supernatant was affinity purified on 3C10-Sepharose as described above for humane urine derived CD14. The specificity of 3C10 has been mapped to residues 7-10 of human CD14 (J.Biol. Chem. 270:361,1995). These residues are highly conserved in bovine and murine CD14.

The left panel of Figure 18 illustrates a comparative silver staining analysis of 1  $\mu$ g each of affinity purified human urine CD14 (nHu), colostral bovine CD14 (nBo), and OKT3 derived mouse CD14 (nMo). The right panel of Figure 18 illustrates a comparative immunoblot analysis of 250 ng of each of the same three species of CD14, probed with mAb 3C10. As illustrated, the purity of all three preparations was comparable, as was their reactivity with mAb 3C10.

Apparent discrepancies in the molecular weights of CD14 of the three species with the differences in the number of amino acids encoded by their respective cDNAs could be due to co- and/or post-translational modification. In this context, it is apparent that mouse, human, and bovine CD14s contain five, four and three potential N-glycosylation sites respectively.

The capacity of these three CD14 preparations to stimulate murine B cell growth was assessed. As illustrated in Figure 19, CD14 isolated from the three species had comparable specific activity, and was active in the ng/ml concentration range. The results also demonstrate that isogenic material is functional, specifically, murine CD14 can stimulate murine B cells. The results also demonstrate that colostral CD14 is not peculiar in its capacity to stimulate B cells, and thus it is not likely a special form of the molecule which is generated in the specialized circumstances of lactation.

# Growth Promoting Activity of nBo-LAIT on Human Cord Blood B Cells

Having observed a variety of bioactivities of nBo-LAIT on murine B cells, effects on the physiology of human B cells were examined. Two sources of B cells were utilized. Since one possible role of LAIT-protein is involvement in potentiating the development of the neonatal immune system, its capacity to stimulate the growth of B cells derived from the neonate, specifically, those isolated from cord blood was assessed.

Cord blood was diluted 1:1 in phosphate buffered saline (PBS), and overlayed onto Percoll (Pharmacia),  $\rho$  = 1.077. The gradient was centrifuged as described in connection with Figure 1A. The  $\rho$  = 1.077/1.000 interface was harvested and washed twice in PBS supplemented with 5% fetal bovine serum (FBS). The resulting leukocyte preparation was then stained with fluorescein conjugated mAb specific for the B cell membrane marker CD72. CD72 positive cord leukocytes were then positively selected by FACS, resulting in purities of

>98%. These positively selected B cells were then cultured in serum free defined medium, as for murine B cells. The only difference in the growth assays for murine and human B cells, was that the latter were pulsed with thymidine at 60 hours, for 12 hours, rather than at 40 hours, for 6 six hours.

As illustrated in Figure 20A, nBo-LAIT acts with mAbs specific for IgK and IgA in its capacity to induce the growth of neonatal B cells. While both immobilized (plate bound) anti-light chain mAbs and 2  $\mu$ g/ml nBo-LAIT induce an increase in thymidine uptake over background, individually, the combination of the two supported a further 5-fold increase.

These results indicate that it might be possible that LAIT protein consumed by the breast fed neonate functions as a T cell surrogate in aid of stimulating B cells which have encountered antigen to grow and differentiate into Ig secreting cells, in the absence of a fully developed T cell compartment (J. Exp. Med. 169:2149, 1989; Science 245:749, 1989; Intl. Immunol. 2:859, 1990; Intl. Immunol. 2:869, 1990).

#### CD14 in Human Colostrum and Normal Serum

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To determine the concentration of CD14 present in colostrum and normal serum, colostral samples from two donors, and serum samples from five healthy donors, were assayed for the presence of CD14 using a specific ELISA (IBL-Hamburg). As shown in Table 3, serum from this cohort of healthy individuals contained CD14 concentrations ranging from 1.7-3.2  $\mu$ g/ml, and was gender independent. These values correspond well with those reported by Grunwald et al. (J. Immunol. Method 155:225,1992).

TABLE 3			
DONOR	GENDER	µg/ml*	
A.M.	F	2.8	
N.J.	F	2.7	
M.J.	М	1.7	
E.K.	М	2.9	
A.D.	М	3.2	

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\* 2.6-3.4  $\mu$ g/ml according to ELISA kit manufacturer (IBL, Hamburg)

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As illustrated in Figure 21, the CD14 content in colostrum taken at 22 hours post-partum (A.D.), and early breast milk taken at four days post-partum (S.B.), contained

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roughly 20-fold higher concentrations of CD14 than did normal serum. Multiple samples extending to 78 days post-partum were obtained from one donor (S.B.), and while CD14 concentrations dropped considerably compared to that observed at day 4, they remained roughly 3-5-fold higher than that observed in normal serum, throughout the screening period assayed (Figure 21).

No information regarding the concentration of serum CD14 in lactating women is yet available. Thus, it remains to be determined whether the high concentrations of CD14 observed in colostrum and breast milk are restricted to these fluids, or reflect a generalized increase in CD14 concentration in all body fluids of lactating women.

It may be that the transient exposure of the neonatal immune system to the B cell tropic growth and differentiation activity of colostral CD14 plays a part in development of the neonatal immune response machinery. The physiological relevance of the presence of this activity in colostrum is consistent with the observation that, as described above, T cell function in the neonate is compromised, possibly due to the presence of high concentrations of TGFβ1 and TGFβ2 in colostrum and early breast milk (J. Cell. Biol. 105:1039, 1987; Cell 49:437, 1987; EMBO J. 6:1633, 1987). As shown in Table 2, submitogenic concentrations of CD14 in combination with submitogenic concentrations of mAb specific for membrane immunoglobulin, supports the activation of B cells. CD14 might function as a T cell surrogate within the developing neonatal immune system. As such, a neonate can benefit from the use of CD14 as an infant formula additive by exposure to its immune-stimulating effects absent from synthetic formula.

### Growth Promoting Activity of nBo-LAIT on Human Tonsil B Cells

The bioactivity of nBo-LAIT on B cells isolated from adults was assessed, in isolation, and in combination with immobilized (plate bound) anti-light chain mAbs, to stimulate B cells isolated from human tonsils.

Tonsil B cells were prepared by negative selection. Tonsil leukocytes were prepared as for cord blood leukocytes. The resulting population was labelled with biotinylated mAb specific for CD3 $\epsilon$  (Becton Dickenson), followed by labelling with iron containing "microbeads" (Becton Dickenson). After one wash, the labelled population was passed through a MACS (Becton Dickenson), and the effluent collected. This population contained <1% T cells, and >97% B cells as assessed by immunofluorescence staining with lineage specific mAbs. These B cells were then subjected to further fractionation on Percoll discontinuous density gradients, identical to those used for the isolation of high buoyant density murine B cells. The assays described used those B cells banding at the  $\rho = 1.085/1.079$  interface. These negatively selected, density fractionated resting B cells were cultured as described below, pulsed, and harvested as for cord blood B cells.

As illustrated in Figure 20B, and in contrast to results obtained with B cells

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isolated from neonates, nBo-LAIT, in isolation, present at concentrations as low as 300 ng/ml stimulated robust growth of these resting tonsil B cells. Further, the response at some concentrations of nBo-LAIT was substantially enhanced when assessed in combination with immobilized anti-light chain mAbs (Figure 20B).

Mature human B cells are susceptible to the growth promoting activities of nBo-LAIT, which are amplified in combination with simultaneous ligation of the B cell antigen receptor. These results characterize the potential utilization of LAIT-protein in vaccine vehicles, in aid of increasing their adjuventicity, or by possibly reducing the need for adjuvants.

A limitation of vaccination technology is the immunogenicity of a particular antigen preparation. Certain adjuvants are thought to function by recruiting and activating antigen specific T cells. CD14, as a T cell surrogate for antigen specific B cell responses, may provide an improved means to activate antigen specific B cells such that they will not only expand and differentiate into antibody secreting cells, but would, once activated, function as efficient APC for the recruitment of T cells. This would enhance both the propagation of the specific immune response and T cell mediated isotype switching.

T cell immune deficiencies are known. Immunodeficiency states associated with T cell dysfunction due to the lack of expression of gp39 (CD40L) (which maps to the X chromosome) have been characterized: (i) X-linked hyper IgM syndrome (HIM); (ii) common variable immunodeficiency (CVI); and (iii) X-linked agammaglobulinemia (XLA). In some of these disease states (HIM), T cells isolated from patients have been shown to be unable to activate B cells (Science 259:990, 1993), and this phenotype correlates with the absence of functional gp39 (CD40L). In these circumstances, CD14, either targeted for the induction of specific humoral responses, or administered as a polyclonal B cell activator could function to induce/maintain levels of isogenic Ig consistent with protection against the daily barrage of potential environmental pathogens.

The presence of CD14 in colostrum is consistent with its role in stimulating B cells within the suckling neonate. The effectiveness of CD14 in aiding development of neonate immune systems can be evaluated in an animal model.

CD14 deficient females, created through targeted disruption of the CD14 locus, will be mated with either heterozygous, or CD14 deficient males. This will enable the assessment of the effects of the absence of colostral CD14 on B cell development in pups that do, or do not express CD14. Specifically, B cell ontogeny and the accumulated development of serum IgM and IgG levels will be compared, as well as the capacity of these pups to mount specific immune responses.

Further, the role that serum CD14 (sCD14) plays in the maintenance of circulating levels of "natural" IgM can be assessed. Levels of circulating IgG and IgM are under distinct control. Serum IgG is virtually absent in mice reared in an antigen free environment,

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while IgM levels are unaltered. Towards addressing the potential role of sCD14 in the regulation of serum IgM levels, CD14 sufficient and deficient mice derived from the above matings will be reared gnotobiotically.

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Dysregulated expression of sCD14 is associated with the pathology of specific disease states. The level of sCD14 in the serum of patients with rheumatoid arthritis (RA) is elevated (Clin. Exp. Immunol. 91(2):207, 1993). It has also been reported that there is an increase in the number of activated CD14\* monocytes in the synovium of RA patients (Br. J. Rheumatol. 29(2):84, 1990; J.Rheumatol. 22(4):600, 1995). While it remains to be determined whether the level of sCD14 in synovial fluid of RA patients is elevated, CD14\* monocytes, upon activation, express membrane associated proteases which can cleave membrane CD14, resulting in the production of sCD14 (Eur. J. Immunol. 25:604, 1995). Consistent with the capacity of sCD14 to activate human B cells, described herein, the synovial fluid of RA patients contains high frequencies of activated B cells, at least some of which may be producing rheumatoid factor (Clin. Immuno. Immunopathol. 31(2):272, 1984; Clin. Exp. Immunol. 55(1):91, 1984). Thus, a paradigm emerges, involving the increased production of sCD14 in RA patients, and its possible involvement in the activation of B cells resulting in the production of rheumatoid factor. Antibody mediated clearance of sCD14 may therefore result in the amelioration of symptoms mediated by dysregulation of B cell activation and rheumatoid factor production in RA patients. Further, antibody mediated clearence of sCD14 would ameliorate inflammation supported by sCD14 induction of pro-inflammatory cytokines by monocytes (Eur. J. Immunol. 24:1779, 1994).

Routine production of human monoclonal antibodies (mAbs) has been difficult for a number of reasons, not the least of which is the inability to enrich for activated human B cells of desired antigen specificity. The capacity of sCD14 to induce human B cells of desired antigen specificity. The capacity of sCD14 to induce human B cell growth and differentiation in vitro, affords its possible utilization in the production of antigen specific mAbs. We show herein that sCD14 at high concentrations (0.5-1 µg/ml) activates human B cells in a polyclonal fashion. However, sub-optimal mitogenic concentrations of sCD14 are shown to synergize with mAb specific for the B cell receptor for antigen (BcR). Thus, sub-optimal concentrations of sCD14 preferentially activate those B cells which receive a complementary signal through the BcR. The BcR specific mAb functions as an antigen surrogate in these circumstances. If specificity is imposed on the delivery of the BcR signal, the ensuing B cell response would also be specific. Thus, when anti-BcR is replaced by a specific antigen, the synergistic stimulus provided by the simultaneous presence of sCD14 would be focussed on the antigen specific B cells, exclusively. Thus, the ensuing production of antibody would be antigen specific. Populations of B cells activated in this fashion would be highly enriched for activated, antigen specific B cells, and would therefore facilitate the production of human hybridomas secreting mAb of desired specificity.

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The effectiveness of CD14 as an adjuvant in vaccination technology can be evaluated using an animal model. Bo- and Hu-CD14 will be modified with the hapten TNP. Haptenated material will be assessed for its capacity to induce polyclonal B cell activation in vitro, to insure that haptenation has not altered CD14 bioactivity. Conjugates will be injected subcutaneously, or intramuscularly, and over time, serum will be assessed for its content of specific antibody. Using another series of mice, draining lymph nodes will be collected, and contained antibody secreting cells enumerated. In addition, some recipients will be immunized with mixtures of varying amounts of CD14 and either protein or cellular antigen. Serum antibody titres, as well as antigen specific, and total Ig secreting cells will be enumerated.

Toxicity of CD14 can be evaluated in acute intravenous studies in mice, rats and monkeys. Acute subcutaneous irritation studies in rats can be performed, as well as in the long term, studies involving multiple subcutaneous and intravenous injections in the three species. Gross pathologic and histopathologic assessment will be performed, as well as serum chemistry and hematological analyses. The genotoxic potential can be assessed in mammalian cells in vitro, and in a mouse micronucleus assay. Teratogenic potential can be assessed in pregant mice, rats, and monkeys.

In administering CD14 to a human subject, conventional pharmaceutical practice can be employed. As an additive to infant formula, it might be added to the formula at the time of manufacture. It might be prepared as a tablet or capsule, or powder for mixing just prior to administration. In the case of vaccine preparation, it might be included as part of a vaccine prepared according to otherwise standard procedures. Administration could be by any convenient means, for example, intravenous, subcutaneous, intramuscular, intraventricular, intracapsular, intraspinal, intracisternal, intraperitoneal, or oral administration.

Parenteral formulations may be in the form of liquid solutions or susupensions.

Methods known in the art for making formulations can be found in, for example, "Remington's Pharmaceutical Sciences." Formulations for parenteral administration may, for example, contain as excipients sterile water or saline, polyalkylene glycols such as polyethylene glycol, vegetable oils, hydrogenated naphthalenes, etc.

The concentration of CD14 for administration will vary depending upon, for example, the dosage to administered and the route of administration.

In terms of variation from a native amino acid sequence of CD14, at the very least, conservative substitutions could be made. Conservative substitutions are described in the patent literature, as for example, in United States Patent No. 5,2264,558. It is thus expected, for example, that interchange among non-polar aliphatic neutral amino acids, glycine, alanine, proline, valine and isoleucine, would be possible. Likewise, substitutions among the polar aliphatic neutral amino acids, serine, threonine, methionine, asparagine and glutamine could possibly be made. Substitutions among the charged acidic amino acids, aspartic acid and glutamic acid, could probably be made, as could substitutions among the

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charged basic amino acids, lysine and arginine. Substitutions among the aromatic amino acids, including phenylalanine, histidine, tryptophan and tyrosine would also likely be possible. These sorts of substitutions and interchanges are well known to those skilled in the art. Other substitutions might well be possible. Of course, it would also be expected that the greater the percentage of homology of a variant protein with a naturally occurring protein, the greater the retention of metabolic activity.

### WHAT IS CLAIMED IS:

- 1. A polypeptide capable of activating mammalian B cells having the amino acid sequence SEQ ID NO:4, or fragment of said polypeptide which retains biological activity of said polypeptide; or a variant of said polypeptide which retains biological activity of said polypeptide; a conservatively substituted variant of the polypeptide; or conjugates of the fragment or variant thereof having biological activity of said polypeptide.
- 2. A polypeptide having SEQ ID NO:4 or a conservatively substituted variant thereof.
- 3. A polypeptide of claim 1 or 2 in which the polypeptide is co- and/or post-translationally modified.
- 4. A nucleic acid molecule comprising a nucleotide sequence encoding for a polypeptide having an amino acid sequence according to claims 1 and 2.
  - 5. A nucleic acid molecule comprising a nucleotide sequence encoding for a polypeptide having an amino acid sequence SEQ ID NO:4.
  - 6. A nucleic acid molecule according to claims 4 and 5 which is a cDNA.
- 15 7. A nucleic acid molecule according to claims 4 and 5 which is a genomic DNA sequence.
  - 8. A nucleic acid molecule according to claim 5 which is a cDNA molecule comprising the nucleotide sequence SEQ ID NO:1.
- A method for preparing a polypeptide having sequence SEQ ID NO:4 or a co- and/or post-translationally modified form of said protein from bovine colostral whey comprising
   isolating the polypeptide from the whey.
  - 10. A method of claim 9, wherein isolating the polypeptide includes salting out of proteins from clarified bovine colostral whey.
  - 11. A method of claim 10 wherein further isolation of the polypeptide includes anion exchange chromatography.
- 12. A method of claim 11 wherein further isolation of the polypeptide includes sieving column chromatography.

- 13. A method of claim 12 wh rein further isolation of the polypeptide includes hdroxy apatite column chromatography.
- 14. A method of activating B cells using a polypeptide having the amino acid sequence SEQ ID NO:4, SEQ ID NO:5, or SEQ ID NO:6, or fragment of said polypeptide which retains biological activity of said polypeptide; or a variant of said polypeptide which retains biological activity of said polypeptide; a conservatively substituted variant of the polypeptide; or conjugates of the fragment or variant thereof having biological activity of said polypeptide.
- 15. A method of claim 14 wherein analogues of the polypeptide SEQ ID NO:4, SEQ ID NO:5, or SEQ ID NO:6 derived from substitutions, deletions, or additions of amino acids of said polypeptides providing the three dimensional conformation of said polypeptides is preserved.
  - 16. A method of claim 15 wherein the polypeptide is a conservatively substituted variant of SEQ ID NO:4, SEQ ID NO:5, or SEQ ID NO:6.
- 15 17. A method of claim 14, 15 or 16 in which the polypeptide is co- and/or post-translationally modified.
  - 18. A method of activating B cells in a mammal in need of such activation by administering a polypeptide according to claim 14, 15,16, or 17.
  - 19. A method of claim 18 in which the mammal is a human subject.
- 20 20. A method of claim 19 wherein the human is an infant.
  - 21. A method of claim18 wherein the mammal in need of such activation has CD40 negative or CD40 deficient B cells.
  - 22. A method of claim 21 wherein the mammal is a human subject.
  - 23. A method of claim 22 wherein the human is an infant.
- 25 24. A method of claim 18 wherein the mammal is suffering a T cell immunedeficiency, or an allergy.
  - 25. A method of claim 24 wherein the mammal has CD40L negative or CD40L deficient T cells.



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- 26. A method of claim 25 wherein the mammal is a human subject.
- 27. A method of claim 26 wherein the human is an infant.
- 28. A method of inducing growth and differentiation of B cells to high rate immunoglobulin secreting cells using a polypeptide having the amino acid sequence SEQ ID NO:4, SEQ ID NO:5, or SEQ ID NO:6, or fragment of said polypeptide which retains biological activity of said polypeptide; or a variant of said polypeptide which retains biological activity of said polypeptide; a conservatively substituted variant of the polypeptide; or conjugates of the fragment or variant thereof having biological activity of said polypeptide.
- 29. A method of claim 28 wherein analogues of the polypeptide SEQ ID NO:4, SEQ ID NO:5,
   10 or SEQ ID NO:6 are derived from substitutions, deletions, or additions of amino acids of said polypeptides providing the three dimensional conformation of said polypeptides is preserved.
  - 30. A method of claim 28 wherein the polypeptide is a conservatively substituted variant of SEQ ID NO:4, SEQ ID NO:5, or SEQ ID NO:6.
- 31. A method of claim 28, 29, or 30 in which the polypeptide is co- and/or posttranslationally modified.
  - 32. A method of inducing growth and differentiation of B cells in a mammal in need of such activation by administering a polypeptide according to claim 28, 29, 30, or 31.
  - 33. The method of claim 32 wherein the mammal is a human subject.
  - 34. A method of claim 31 wherein the human is an infant.
- 20 35. A method of preparing a pure protein or co- and/or post-translationally modified form of the protein having amino acid sequence SEQ ID NO:4 including conservatively substituted variants tehreof, comprising of:
  - culturing a prokaryotic or eukaryotic host capable of expressing said protein or co- and/or post-translationally modified form of said protein under culturing conditions;
  - (ii) expressing said protein or co- and/or post-translationally modified form of said protein; and
  - (iii) recovering the said protein from the culture.

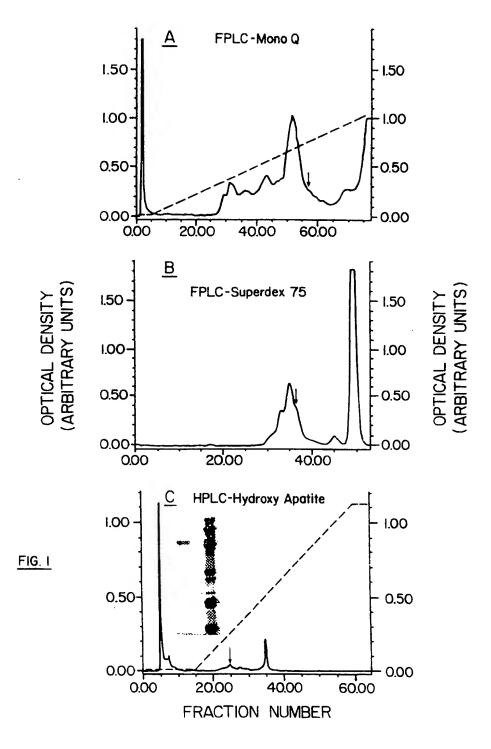


- 36. The use of a polypeptide of claim 14, 15, 16, or 17 for the manufacture of a medicament for activating B cells in a mammal in need of such activation.
- 37. A method of manufacturing infant formula which comprises incorporating a polypeptide of claim 14, 15, 16, or 17 into the formula.
- 5 38. A method of manufacturing a vaccine which comprises incorporating a polypeptide of claim 14, 15, 16, or 17 into the vaccine formula.
  - 39. The method of claim 38, comprising the step of conjugating antigen and polypeptide of claim 14, 15, 16, or 17 into the vaccine formula.
- 40. The method of claim 38, further comprising the step of mixing antigen and polypeptide of claim 14, 15, 16, or 17 into the vaccine formula.
  - 41. A method of vaccinating a patient comprising administering an antigen and a polypeptide of claim 14, 15, 16, or 17 to the patient.
  - 42. The method of claim 41 wherein the antigen and polypeptide of claim 14, 15, 16, or 17 is administered in a single step.
- 15 43. The method of claim 41 wherein the antigen and polypeptide of claim 14, 15, 16, or 17 is administered in separate steps.
  - 44. A kit for the preparation of a vaccination comprising a predetermined amount of an antigen and a predetermined amount of a polypeptide of claim 14, 15, 16, or 17.
- 45. A method of administering to a mammal in need of an antibody raised against a polypeptide with an amino acid sequence of SEQ ID NO:5 or fragment of said polypeptide which retains biological activity of said polypeptide; or a variant of said polypeptide which retains biological activity of said polypeptide; a conservatively substituted variant of the polypeptide capable of binding said polypeptide, fragment or variant thereof and thereby reducing or inhibiting the activity of human B cells that are otherwise hyperactivated as a result of high serum levels of a polypeptide with an amino acid sequence of SEQ ID NO:5 or functional derivatives thereof.
  - A method of claim 45 wherein the mammal is a human subject.

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- 47. A method of enriching for mammalian B cells secreting monoclonal antibody of desired antigen specificity comrising of activating said B cells according to claim 14, 15, 16, or 17 at sub-optimal levels in concert with an antigen to which antibodies are to be raised against.
- 48. A method according to claim 47 in which the mammalian B cells are of human origin.
- 5 49. A method according to claim 47 in which the mammalian B cells are of murine origin.



**SUBSTITUTE SHEET (RULE 26)** 

- 1
  MERASCLLLL LLPLVHVSAT TPEPCELDDE DFRCVCNFSE PQPDWSEAFQ
  D -?--?--NN? ??-?-?
- 51
  CVSAVEVEIH AGGENCEPFL KRVDADADPR QYADTVKALR VRRLTVGAAQ
  V---?-S ---LSL?
- 101 VPAQLLVGAL RVLAYSRLKE LTLEDLKITG TMPPLPLEAT GLALSSLRLR
- NVSWATGRSW LAELOQWLKP GLKVLSIAQA HSPAFSYEQV RAFPALTSLD
- 201 LSDNPGLGER GLMAALCPHK FPAIQNLALR NTGMETPTGV CAALAAAGVO ---S-P A----LR--
- PHSLDLSHNS LRATVNPSAP ROMWSSALNS LNLSFAGLEQ VPKGLPAKLR -Q----? -?V?
- 301 VLDLSCNRLN RAPQPDELPE VDNLTLDGNP FLVPGTALPH EGSM $\underline{NSGVVP}$  L----V
- 351
  ACARSTLSVG VSGTLVLLQG ARGFA
  ???--

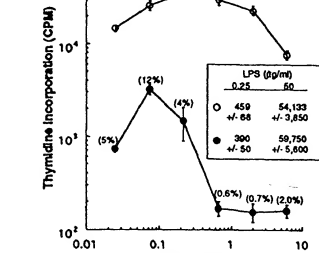


FIG. 3

Untreated 95°C 10'

10





0.1

LAIT (µg/ml)

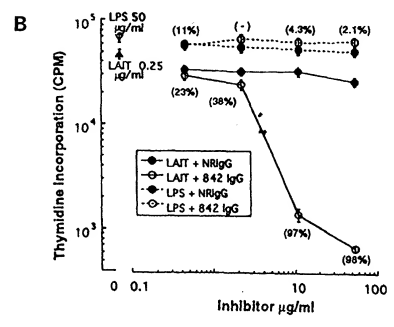


FIG. 4

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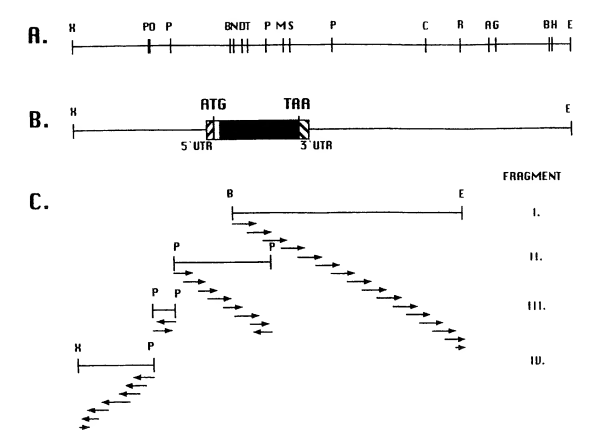


FIG. 5

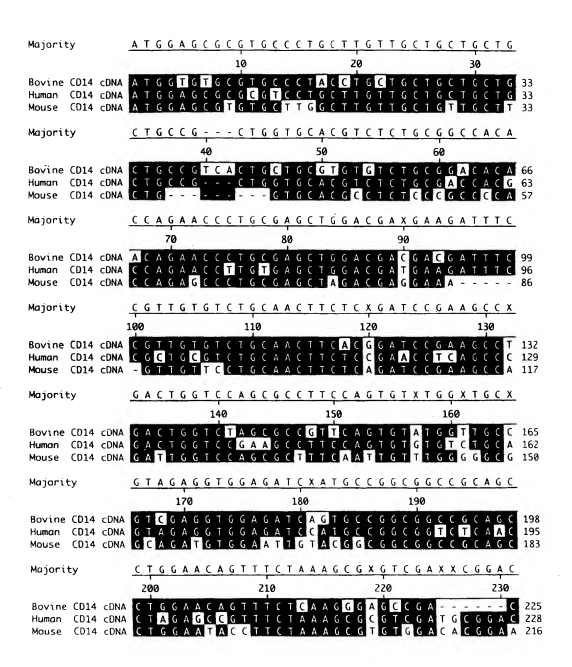


FIG. 6 A



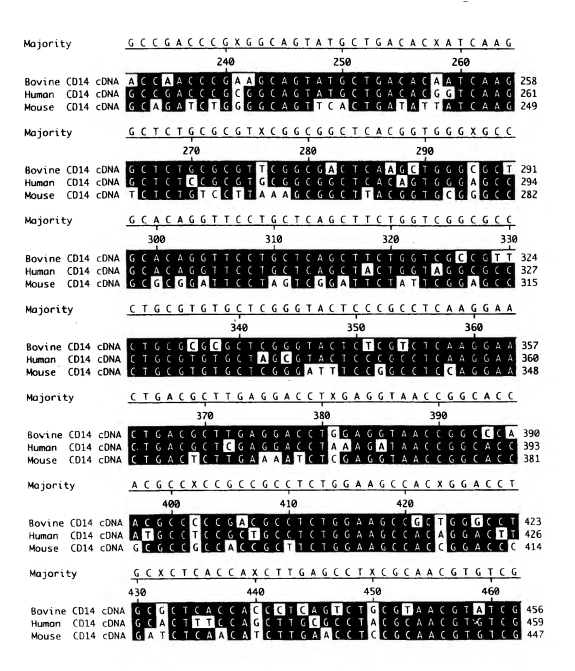


FIG. 6 B



Majority	TGGGCAACAGGGG	тосстоостсос	CGAACTG
	470	480	490
Human CD14 cDNA	T G G A C A A C A G G A G G T G G G C G A C A G G G C G T G G G C A A C A A G G G A	TICTICACICAC	CGAGCTG 492
Majority	CAGCAGTGGCTCAA	GCCTGGXCTCAA	GGTACTG
	500 5	10 520	
Human CD14 cDNA	C A G C A G T G G C T C A A C C A G C A G T G G C T C A A C C A G T G G C T A A A C C A G	GCCAGGCCTCAA	G G T A C T G 525
Majority	AGCATTGCCCAAGC	ACACTCGCTTGC	CTTTTCC
	530 540	550	560
	AGCATTGCCCAAGC	A C A C T C G C T T G C A C A C T C G C C T G C A C A C T C A C T C A A	C T T T T C C 558
Majority	TGCGAACAGGTCCG	CGCCTTCCCGGC	CCTCACC
	570	580	590
Bovine CD14 cDNA Human CD14 cDNA Mouse CD14 cDNA	T G C G C A G G G C T C T C G T G C G A A C A G G T T C G G T G C G A A C A G G T C C G G	CGCCTTCCCGGC	( ( T T A ( ( 591
Majority	ACCCTAGACCTGTC	TGACAATCCTGG	ACTGGGC
	600	610 620	9
Human CD14 cDNA	A C C C T A G A C C T G T C T A'G C C T A G A C C T G T C T A C C T T A G A C C T G T C T	IGACAATCCTGG	A C T G G G C 624
Majority	GAXACGXGGACTGA	TGGCAGCTCTCT	GTCCCCA
	630 640	650	660
Bovine CD14 cDNA Human CD14 cDNA Mouse CD14 cDNA	G A C A C G - G G G C T G A - G A G A C T A - G A G A C T A - G A G A C T A - G A G A C T A - G A G A C T A - G A G A C T A - G A -	TGGCGGCTCTCT	G T C C G A A 653 G T C C C C A 656 G T C C C C T 644
Majority	CAAGTTCCCGGCCC	TCCAAXATCTAG	СССТСС
	670	680	690
Bovine CD14 cDNA Human CD14 cDNA Mouse CD14 cDNA	CAAGTTCCCGGCCA	T C C A A T A T C T A G T C C A G A A T C T A G T C C A A G T T T T A G	C G C T G C G 689

FIG. 6 C

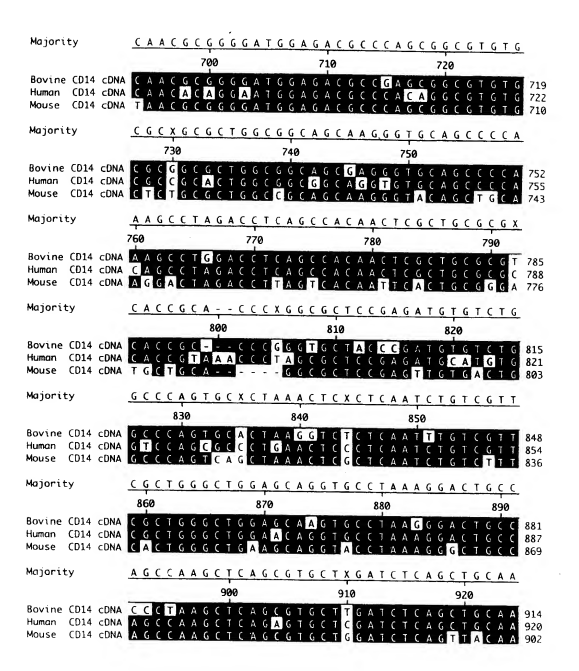


FIG. 6 D



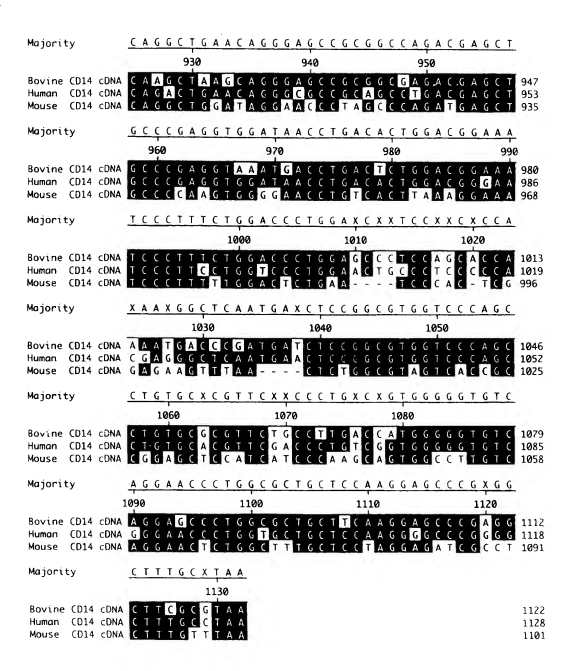


FIG. 6 E



Majority	MERVXXLLLLL	LP-LVHVSAXTP	EPCELDDEDFRC	
	10	20	30	
Bovine CD14 Human CD14 Mouse CD14	MERASCLLLLL	L P S L L R V S A D T T L P - L V H V S A T T P L ~ V H A S P A P P	EPCELDDEDFRC	35 34 30
Majority	VCNFSDPKPDW	SSAFQCXXAVEV	EIXAGGRSLEXF	
	40	50	60 70	
Bovine CD14 Human CD14 Mouse CD14	IA C M E 2 F B O B D M	S	E T ULA C C TEMPS . C S C .	70 59 55
Majority	LKRVDADADPX	QYADTIKALRVR	RLTVGAAQVPAQ	
	80	90	100	
Bovine CD14 Human CD14 Mouse CD14	LKRVDADADPR	Q Y A D T I K A L R V R Q Y A D T V K A L R V R Q F T D I I K S L S L K	PITYCALOUGIC	103 104 100
Majority	LLVGALRVLGY	SRLKELTLEDLE	VTGTXPPXPLEA	
	110	120	130 140	
Bovine CD14 Human CD14 Mouse CD14	L	SRIKELTLEDLE SRIKELTLEDL <mark>K</mark> S <b>G</b> L <b>Q</b> ELTLE <b>N</b> LE	TTCTUODIC	.38 .39 .35
Majority	TGPALXXLXLR	NVSWATGXAWLA	ELQQWLKPGLKV	
	150	160	170	
Human CD14 Mouse CD14	UGLAUSSLRIR	N V S W T T G G A W L G N V S W A T G R S W L A N V S W A T R D A W L A	E L Q Q W L K P G L K V 1	.73 .74 .7 <b>0</b>
Majority	LSIAQAHSLAF	SCEQVRXFPALT	TLDLSDNPXLGE	
	180	190	200 210	
Human CD14	LSIAQAHSPAF	PCAGLSTFEALT SYEQVRAFPALT SCEQVRVFPALS	SLDLSDNPGLGEZ	08 109 105
Majority	RGLMAALCPXK	FPALQXLALRNA	GMETPSGVCAAL	
	220	230	240	
Human CD14		FPAIQNLALRNT	GMETPTGVCAALZ	43 44 240

FIG. 7 A



Majority		<u>A</u>	<u>A</u>	<u>A</u>	R	Ÿ	Q	Р.	Q	<u>S</u>	Ĺ	D	L	<u>S</u>	<u>H</u>	N	<u>S</u>	L	R	X	T	-	Α	P	G	A	P	R	C	X	W	Р	S	Α	L	N	
					ā	250	9									:60										270										280	,
	D14 D14 D14	A	A	A	G	V	ò	PPL	Ĥ	S S G	L	D D	Ł	S	Н Н	Ν	5	Į.	R R R	V A D	I A	- V -	A N A	Р Р -	G G	Λ	T P	D	c	M	W	P S P	S S	A		RNN	277 279 273
Majority		S	L	N	L	S	F	A	G	L	Ε	Q	v	Р	ĸ	G	L	Ρ	Α	ĸ	L	5	٧	L	D	L	S	c	N	R	L	х	R	X	P	х	
										2	290	)								3	00	)									310						
Bovine CD Human CD Mouse CD	D14 D14 D14	5	1	Ν	1	5	F	Α	G	T.	F	0	V	P	К	6	1	C)	Δ	K		D	V	1	0		<	c	м	б	L L	SND			P P	٦.	312 314 308
Majority		Р	D	E	L	Р	Ε	٧	х	N	L	T	L	D	G	N	Р	F	L	D	Р	G	Х	X	x	X	н	X	x	X	м	N	s	G	٧	v	
					:	320	)								3	30	)								3	346	)								3	50	
	014 014 014	P	D	Ε	L	Ρ	EQ	٧	N D G	N	L	T	L	D	G	N	P	F		V	Р	G	T	Α	Ĺ	P	m	F	G	ς	М	M	S S S	G G	V V V	> > >	347 349 340
Majority		Р	A	C	A	R	5	X	L	х	٧	G	٧	S	G	T	L	A	L	L	Q	G	A	R	G	F	Α										
										3	366	)								3	70	)															
	014 014 014	PPT		į					Q	S	٧	GA	٧	Š	G G		L	A V A	L L L		Q	GGG		R	COL		_										373 375 366

FIG. 7 B

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# Α

# Baculovirus expression system

Forward 5'- GCT AGC GCT AGC CAC CAT GGT GTG CGT GCC CTA CCT GCT - 3'

Reverse 5' - GCT AGC GCT AGC CGC GAA GCC TCG GGC TCC TTG AAG - 3'

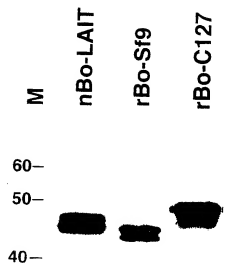
# В

# Mammalian expression system

Forward 5' - CTC GAG CTC GAG GCT AGC CAC CAT GGT GTG CGT GCC - 3'

Reverse 5' - CTC GAGCTGAG GGA TCC CTA AGC GTA ATC TGG AAC - 3'

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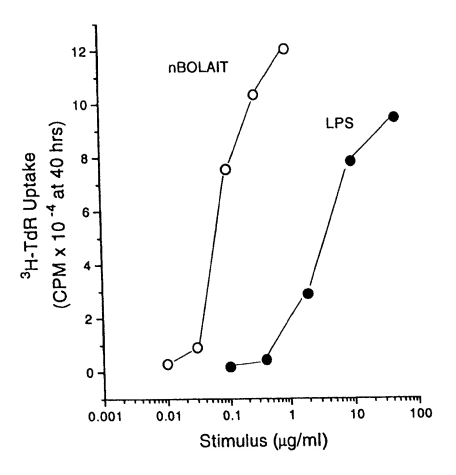


FIG. 10

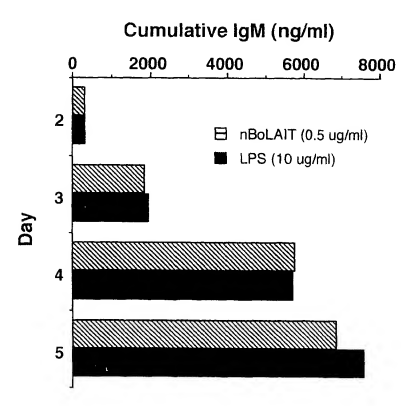


FIG. 11

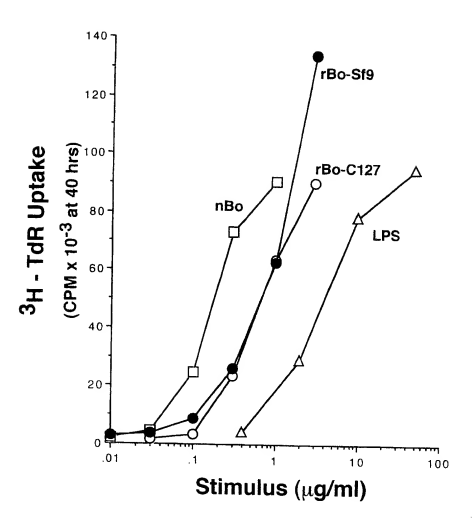


FIG. 12

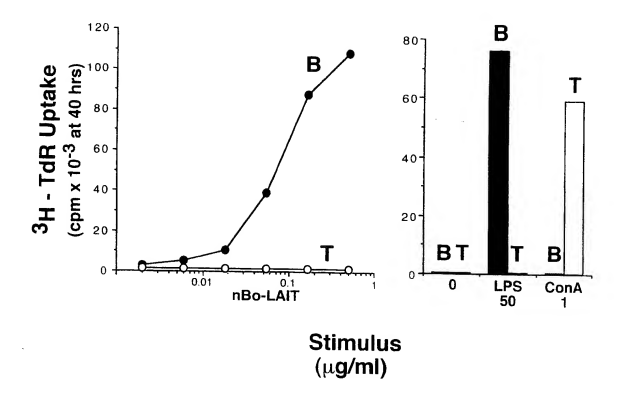
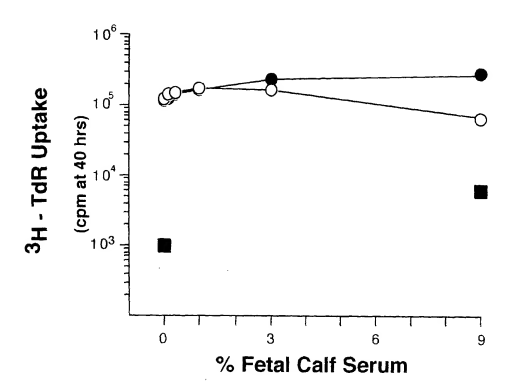
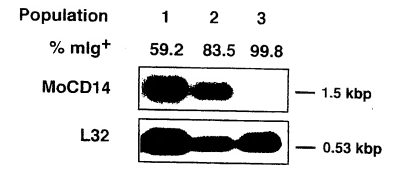


FIG. 13



- unstimulated
- O nBo-LAIT 0.5µg/ml
- LPS 50 μg/ml

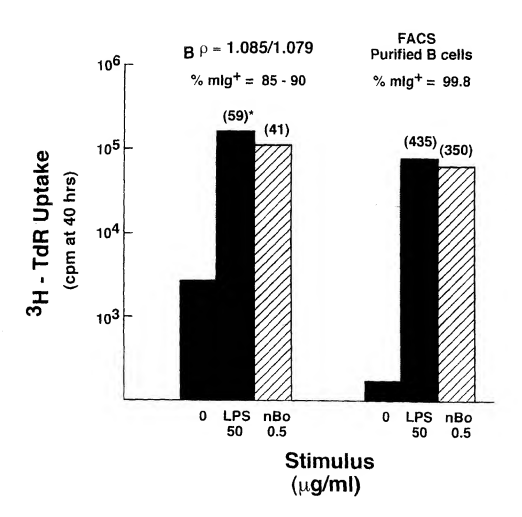
FIG. 14



Populations: 1 - unfractionated splenocytes

2 - T-depleted,  $\rho$ =1.085/1.079 splenocytes

3 - FACS purified mlg<sup>+</sup> B cells from population 2



\* ( ) = index of stimulation

FIG. 16

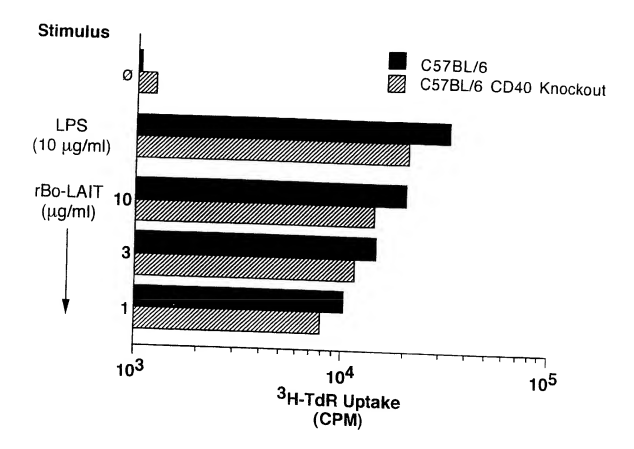
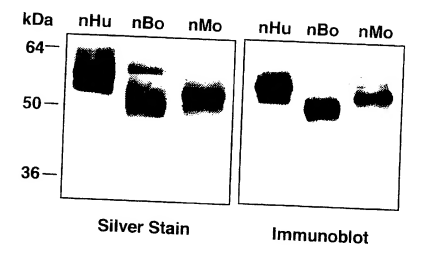


FIG. 17



# **Derivation**

nHu - urine from nephrotic patient

nBo - colostrum

nMo - OKT3-hybridoma supernatant